Calsyntenin-1 shelters APP from proteolytic processing during anterograde axonal transport

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Summary
Endocytosis of amyloid-β precursor protein (APP) is thought to represent the major source of substrate for the production of the amyloidogenic Aβ peptide by the β-secretase BACE1. The irreversible nature of proteolytic cleavage implies the existence of an efficient replenishment route for APP from its sites of synthesis to the cell surface. We recently found that APP exits the trans-Golgi network in intimate association with calsyntenin-1, a transmembrane cargo-docking protein for Kinesin-1-mediated vesicular transport. Here we characterized the function of calsyntenin-1 in neuronal APP transport using selective immunosolation of intracellular trafficking organelles, immunocytochemistry, live-imaging, and RNAi. We found that APP is co-transported with calsyntenin-1 along axons to early endosomes in the central region of growth cones in carriers that exclude the α-secretase ADAM10. Intriguingly, calsyntenin-1/APP organelles contained BACE1, suggesting premature cleavage of APP along its anterograde path. However, we found that APP contained in calsyntenin-1/APP organelles was stable. We further analyzed vesicular trafficking of APP in cultured hippocampal neurons, in which calsyntenin-1 was reduced by RNAi. We found a markedly increased co-localization of APP and ADAM10 in axons and growth cones, along with increased proteolytic processing of APP and Aβ secretion in these neurons. This suggested that the reduced capacity for calsyntenin-1-dependent APP transport resulted in mis-sorting of APP into additional axonal carriers and, therefore, the premature encounter of unprotected APP with its ectodomain proteases. In combination, our results characterize calsyntenin-1/APP organelles as carriers for sheltered anterograde axonal transport of APP.

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Introduction
Excessive production of pathogenic amyloid-β (Aβ) peptide from amyloid-β precursor protein (APP) is considered as the biochemical hallmark of Alzheimer’s disease (Haass and Selkoe, 2007). Aβ formation requires proteolytic cleavage of APP in the juxta membraneous region of its ectodomain by the β-secretase BACE1, followed by intramembrane cleavage by γ-secretase. Competing ectodomain cleavage within the Aβ segment of APP by α-secretase, followed by intramembrane cleavage by γ-secretase, produces the non-amyloidogenic p3 peptide and, thereby, precludes Aβ formation (Sisodia, 1992a). APP has been found at the plasma membrane and, for a fraction of it, internalization through endocytosis and recycling back to the cell surface have been reported (Haass et al., 1992; Koo et al., 1996; Yamazaki et al., 1996; Marquez-Sterling et al., 1997; Groemer et al., 2011). Along its recycling itinerary, APP may be cleaved by α-secretase at the plasma membrane (Sisodia, 1992b; Kojro and Fahrenholz, 2005) and by β-secretase in the early endosome (EE) (Koo and Squazzo, 1994; Perez et al., 1999; Grbovic et al., 2003; Kinoshita et al., 2003; Carey et al., 2005; Rajendran et al., 2006; Small and Gandy, 2006; He et al., 2007).

Based on these findings it is now widely accepted that EEs are a major site of β-secretase activity and that APP internalized from the plasma membrane plays an essential role in Aβ generation. Accordingly, targeted inhibition of endosomal BACE1 through a transition-state inhibitor linked to a sterol was recently shown to reduce Aβ production in vitro and in vivo (Rajendran et al., 2008).

In neurons, the presynaptic nerve is a major site for the release of the soluble ectodomain of APP after proteolytic cleavage by α-secretase (sAPPα) (Nitsch et al., 1992). The irreversible nature of proteolytic processing requires continuous replenishment of APP by anterograde axonal transport. Already two decades ago kinesin was identified as the main molecular motor required for the transport of APP-containing vesicles (Koo et al., 1990; Ferreira et al., 1993; Kaether et al., 2000). Yet, the molecular components and interactions that mediate the connection between APP-containing transport vesicles and Kinesin-1 motors are still controversial. A direct, high-affinity interaction between the cytoplasmic segment of APP and the tetratricopeptide repeat region of the light chains of Kinesin-1 was proposed based on immunoprecipitations (Kamal et al., 2000). An indirect
connection between APP and Kinesin-1 motors was proposed based on reports that c-Jun N-terminal kinase-interacting protein 1 (JIP1b) would act as a bridging protein by simultaneously binding both APP and kinesin light chain-1 (Matsuda et al., 2003; Inomata et al., 2003). However, both mechanisms were questioned by subsequent work by several independent labs that failed to reproduce the direct binding between APP and Kinesin-1 (Lazarov et al., 2005) and the fact that absence of JIP1b did not affect transport of APP (Kins et al., 2006). These objections were backed by a series of reports indicating that APP variants lacking the cytoplasmic segment were still efficiently delivered to axons in several distinct model systems of axonal transport (Tienari et al., 1996; Torroja et al., 1999; Rusu et al., 2007; Back et al., 2007; Szodorai et al., 2009). These results, together with the recent observation that distinct forms of APP are independently transported in separate carrier vesicles (Muresan et al., 2009), suggest that further scrutiny is required to resolve the detailed mechanism(s) of anterograde axonal transport of APP.

We recently found that APP exits the trans-Golgi network (TGN) in intimate association with calsyntenin-1, a transmembrane cargo-docking protein for Kinesin-1-mediated anterograde axonal transport of membrane-bounded organelles (Vogt et al., 2001; Hintsch et al., 2002; Konecna et al., 2006; Ludwig et al., 2009). Like APP, calsyntenin-1 is subject to two-step proteolytic processing. After the first cleavage in the juxtamembrane region of its ectodomain by ADAM10 and ADAM17, but not the β-secretase BACE1, are capable of cleaving calsyntenin-1’s ectodomain (Hata et al., 2009).

Using organelle immunoisolation and proteomics, we recently demonstrated that calsyntenin-1 organelles contain components characteristic of vesicles of endosomal pathways (Steuble et al., 2010). Axons contained at least two distinct, non-overlapping calsyntenin-1-containing transport packages, one characterized by the presence of APP and early-endosomal markers (Rab5), the other with recycling-endosomal markers (Rab11) and no APP (Steuble et al., 2010). In accordance with the identification of calsyntenin-1(APP vesicles as a distinctive carrier for anterograde axonal transport of APP we found that calsyntenin-1 and APP exit the TGN together in a vesicle with early-endosomal characteristics (Ludwig et al., 2009) and that all calsyntenin-1 vesicles are transported anterogradely along axons (Konecna et al., 2006). RNAi studies in cultured neurons indicated that down-regulation of calsyntenin-1 results in enhanced APP processing (Ludwig et al., 2009; Vagnoni et al., 2012). Based on the observation of reduced levels of calsyntenin-1 in brains of humans affected with Alzheimer’s disease a pathogenic role of calsyntenin-1 dysfunction and Alzheimer’s disease was suggested (Vagnoni et al., 2012).

Here we set out to characterize the role of calsyntenin-1 and calsyntenin-1-containing vesicles in axonal APP transport and proteolytic processing. Because work of several laboratories located proteolytic processing of APP by α- and β-secretases in the axonal periphery, we wondered whether anterograde axonal transport of full-length APP included mechanisms to protect APP from its processing proteases during transport to its peripheral destination. The β-secretase has been unequivocally identified as the aspartyl protease BACE1 over a decade ago (Vassar, 2004). In contrast, the molecular identity of α-secretase has long remained controversial. The most frequently discussed candidates were members of the ADAM (a disintegrin and metalloprotease) family, ADAM9, ADAM10, and ADAM17 (Koike et al., 1999; Lammich et al., 1999; Slack et al., 2001). However, studies over the past decade have accumulated evidence that the physiologically most relevant α-secretase responsible for APP processing in the brain is ADAM10 (Kuhn et al., 2010; Jorissen et al., 2010). The role of ADAM9 in α-secretase cleavage of APP was discredited mainly by its failure to cleave an APP-derived peptide at the α-cleavage site in vitro (Roghani et al., 1999) and the finding that α-secretase-derived cleavage products of APP were not reduced in mice lacking ADAM9 (Weskamp et al., 2002). ADAM17 was shown to be mainly expressed in astrocytes and endothelial cells, but not in neurons (Goddard et al., 2001) and, in accordance with this observation, we did not find ADAM17 in immunisolated calsyntenin-1 vesicles (supplementary material Fig. S1).

We show here that calsyntenin-1(APP organelles that are cotransported anterogradely along axons contain BACE1, but not ADAM10, both at the level of the growth cone and along the axon. The presence of BACE1 raised speculations about a premature cleavage of APP along its anterograde axonal path. However, incubation studies with immunisolated vesicles indicated that APP contained in calsyntenin-1(APP organelles was stable, implying that calsyntenin-1 provides a protective mechanism for axonal transport of APP. After arrival in endosomes of the growth cone, APP may be released from its protective complex via cleavage of calsyntenin-1 by endocyted ADAM10.

**Results**

**BACE1, but not ADAM10, is present in calsyntenin-1/APP organelles**

Full-length APP that is lost by proteolytic cleavage during its local recycling in the growth cone may be replenished by calsyntenin-1-dependent anterograde delivery of APP to EEs of growth cones (Steuble et al., 2010). To further characterize the axonal calsyntenin-1(APP carrier, we immunisolated calsyntenin-1(APP, APP, syntaxin13, Rab11, and Rab5 vesicles and tested them for the presence of ADAM10 and BACE1 (Fig. 1A–E). Conversely, we immunisolated ADAM10 and BACE1 organelles with antibodies directed against cytoplasmic epitopes of ADAM10 and BACE1, respectively, and tested them for their content of calsyntenin-1, APP, as well as a series of established organelle markers (Fig. 1F,G). Organelle immunisolations were performed with specific antibodies against marker proteins and magnetic Dynabeads M-280 decorated with anti-IgG (Fig. 1A). Protocol and specificity of the previously established immunolocalizations of calsyntenin-1, syntaxin13, Rab11, and Rab5 organelles were reported previously (Steuble et al., 2010). The specificity of the immunolocalizations of APP, ADAM10, and BACE1 organelles is demonstrated in Fig. 1B. Calsyntenin-1 immunolocalizations contained BACE1, but not ADAM10 (Fig. 1C, boxed). In contrast, APP, syntaxin13, and Rab5 immunolocalizations contained both BACE1 and ADAM10 (Fig. 1D,E, boxed), while Rab11 immunolocalizations were devoid of both (Fig. 1D, boxed). Because Rab11 immunolocalizations did not contain BACE1, we concluded that BACE1 was selectively associated with early-endosomal calsyntenin-1(APP carriers. The absence of ADAM10 from calsyntenin-1 immunolocalizations indicated separate...
anterograde axonal transport for ADAM10 and calsyntenin-1-associated APP.

Separate trafficking of ADAM10 and calsyntenin-1-associated APP was corroborated by the absence of calsyntenin-1 from ADAM10 immunoisolates (Fig. 1F, black box). Because ADAM10 immunoisolates did not contain Rab11 (Fig. 1F) and Rab11 immunoisolates did not contain ADAM10 (Fig. 1D, boxed), we concluded that ADAM10 was also absent from early endosomes (REs) of the slow recycling route. Together, these results indicated that ADAM10 is transported by a calsyntenin-1-free organelle, presumably on a secretory route.

**ADAM10 is a resident of early endosomes**

Besides the expected plasma-membrane marker syntaxin4 (Fig. 1F, green box), ADAM10 immunoisolates also contained the EE marker Rab5 and the pan-endosomal marker syntaxin13.

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**Fig. 1.** See next page for legend.
BACE1 colocalizes with APP in anterogradely transported axonal calsyntenin-1/APP organelles

BACE1 immunoisolates contained calsyntenin-1, APP, the pan-endosomal marker syntaxin13, and the RE marker Rip11. We used an antibody against Rip11, a Rab11-binding protein, as a substitute marker for Rab11, because we could not find a suitable anti-Rab11 antibody for immunocytochemistry. Axons were distinguished from dendrites by staining for hypo-phosphorylated Tau-1 antibody for immunocytochemistry. Axons were distinguished from dendrites by staining for hypo-phosphorylated Tau-1 (Kempf et al., 1996). As documented in Fig. 2, we found a strong colocalization of BACE1 with calsyntenin-1, APP, and syntaxin13 both along axons and in the C-domain of growth cones, while considerably less overlap was found in the P-domain of growth cones (Fig. 2A, a1–c4; Fig. 2B, a1–c4; Fig. 2C–E). The fractions of Rip11 vesicles containing BACE1 were low (2–5%) in all regions examined (Fig. 2A, d1–d4; Fig. 2B, d1–d4; Fig. 2C–E).

The reduction of BACE1 from calsyntenin-1, APP, and syntaxin13 puncta at the transition from the C- to the P-region is consistent with the notion that axonally transported BACE1 accumulates in EEs and that EEs are the main site of BACE1 action (Koo and Squazzo, 1994; Perez et al., 1999; Kinoshita et al., 2003; Rajendran et al., 2006; Rajendran et al., 2008; He et al., 2007). In summary, the distribution pattern of BACE1 along the axon and in the growth cones strongly resembled that of early-endosomal calsyntenin-1/APP organelles. In contrast, only very little colocalization of BACE1 with Rip11 was found.

To corroborate the association of BACE1 with the anterograde trajectory of axonal calsyntenin-1/APP/BACE1 vesicles we expressed combinations of fluorescently tagged calsyntenin-1, APP, BACE1 in cultured hippocampal neurons and tracked their vesicular migration along axons by live-imaging. Anterograde tracks clearly outnumbered retrograde tracks for all tested components and the tagged molecules co-localized in most of the tracks (Fig. 3A–C). Approximately 73% of calsyntenin-1-eGFP, 69% of calsyntenin-1-mRFP, 74% of APP-mRFP, and 72% of BACE1-eGFP transport packages moved in anterograde direction. Likewise, 77% of calsyntenin-1/APP, 69% of calsyntenin-1/BACE, and 69% of APP/BACE vesicles were associated with anterograde tracks (Fig. 3D–F). These results indicated that a substantial fraction of APP and its β-secretase BACE1 are co-transported anterogradely along axons in calsyntenin-1/APP-positive carriers to EEs in the C-domain of growth cones.

Calsyntenin-1 of the anterograde axonal calsyntenin-1/APP route is proteolytically degraded in early endosomes in the C-region of the growth cone

We previously speculated that early-endosomal, APP-positive and Rab11-negative calsyntenin-1 transport packages cease their anterograde axonal trajectory in EEs of growth cones through proteolytic cleavage of calsyntenin-1 (Steuble et al., 2010). To test this hypothesis, we analyzed syntaxin13, APP, and Rab11 immunoisolates for calsyntenin-1 cleavage products before and after incubation at 37°C for 30 min under conditions that preserve their luminal pH and analyzed calsyntenin-1 processing using Western blotting (Fig. 4A–G). Calsyntenin-1 remained unaltered during incubation of APP immunoisolates (Fig. 4A). The C-terminal fragment (CTF) of calsyntenin-1, which was abundant in APP immunoisolates, was also stable during incubation. Likewise, incubation of Rab11 immunoisolates did not result in calsyntenin-1 proteolysis (Fig. 4B). In contrast, incubation of syntaxin13 immunoisolates resulted in the appearance of cleaved calsyntenin-1 ectodomain (Fig. 4C–F). Concomitantly, the amount of calsyntenin-1 CTF was reduced after incubation.
Fig. 2. See next page for legend.
Evidence for protection of APP from BACE1 during anterograde axonal co-transport in calsyntenin-1 organelles

In contrast to calsyntenin-1, the full-length forms of APP remained unaltered during incubation of syntaxin13 vesicles (Fig. 4C,D,H–K). To understand the band pattern found in the various vesicle immunoisolations we also analyzed the maturational glycosylation pattern by cleaving glycoproteins of the V1 fraction and syntaxin13 immunoisolates with PNGaseF and EndoH (Fig. 4L). The single major band of calsyntenin-1 responded to PNGaseF with a clear down-shift, while EndoH did not induce a change, indicating that calsyntenin-1 was exclusively represented by its mature, fully glycosylated form in these samples. Likewise, all three major bands of APP responded to PNGaseF with a down-shift. EndoH, in contrast, left the upper two bands of APP unchanged, but induced a down-shift of the lowest band. In all samples, we found no differences in the band patterns tested with antibodies against N- and C-terminally located epitopes of APP. In accordance with previous analyses of brain tissue samples (Buxbaum et al., 1998) and of cell lysates from cultured cortical neurons (Hoey et al., 2009), we concluded that the two upper high-molecular mass bands of APP identified in the V1 membrane input and in syntaxin13 immunoisolates (Fig. 4C,D) correspond to glycosylation variants of mature full-length APP, while the lower band represents immature full-length APP carrying high-mannose carbohydrate chains. APP in these samples was not cleaved, as the band patterns obtained with antibodies against the N- and the C-terminus of APP were identical. Because the band pattern of APP in syntaxin13 immunoisolates did not change upon vesicle incubation, we concluded that the bulk of APP is stable in syntaxin13-positive endosomal compartments, including EEs, fast recycling endosomes, as well as the calsyntenin-1-dependent anterograde axonal carriers. The fact that the APP fraction that is degraded in EEs during its peripheral recycling pathway was not detected in syntaxin13 immunoisolates indicates that APP of the calsyntenin-1/APP trafficking route vastly predominates over the amount of APP contained in the other APP-containing vesicles, including early-endosomal APP.

In accordance with the notion of a calsyntenin-1-dependent, protected route for anterograde axonal transport of APP, calsyntenin-1 immunoisolates contained predominantly full-length APP and their incubation did not result in APP processing (Fig. 4M). This is in striking contrast to early-endosomal Rab5 and Rab4 immunoisolates in which cleaved APP was predominant (Steuble et al., 2010). Taken together, these results indicate that APP is not cleaved along its anterograde transport in which it is accompanied by calsyntenin-1. Degradation of APP takes place upon arrival in EEs of growth cones and proteolytic degradation of calsyntenin-1.
Down-regulation of calsyntenin-1 results in premature encounter of APP and ADAM10

Because ADAM10 was excluded from calsyntenin-1/APP vesicles, we assumed that APP and ADAM10 did not encounter each other until both had reached the plasma membrane via distinct pathways. We tested whether reduced calsyntenin-1 levels resulted in altered APP trafficking and premature contact of APP with ADAM10. To this end we down-regulated calsyntenin-1 in hippocampal neurons by RNAi and performed immunocytochemical double-staining for ADAM10 and APP (Fig. 5). The specificity of anti-calsyntenin-1, anti-APP, and anti-ADAM10 antibodies was demonstrated by the absence of an immunofluorescence signal after pre-incubation of the antibody with the corresponding antigen (supplementary material Fig. S2).
The quantification of the band intensities in Western blots (Fig. 6A) showed a 52% down-regulation of calsyntenin-1 (Fig. 5A,B) leading to a substantial increase in the colocalization of APP and ADAM10 both in axons and growth cones (Fig. 5C). Calsyntenin-1-specific shRNA increased the fraction of ADAM10 vesicles containing APP from 6% to 21% in axons and from 9% to 26% in growth cones, compared to treatment with nonsense shRNA (Fig. 5D). Conversely, the fraction of APP vesicles containing ADAM10 increased from 4% to 19% in axons and from 8% to 27% in growth cones in the presence of calsyntenin-1-specific shRNA (Fig. 5E). These results suggest that the reduction in the transport capacity for APP in the sheltered calsyntenin-1/APP pathway results in mis-sorting of APP in the TGN and ectopic TGN exit with other post-Golgi carriers. Mis-sorting leads to premature exposure of APP to ADAM10.

Abrogation of anterograde axonal calsyntenin/APP transport increases both α- and β-cleavage of APP and results in increased Aβ release

We next examined whether mis-sorting of APP into ADAM10-containing carriers affected its proteolytic processing. Upon down-regulation of calsyntenin-1 by RNAi, APP cleavage at both the α- and β-site was enhanced (Fig. 6A,B). The normalized α-C99/β-C83 ratio after calsyntenin-1 RNAi was 1.05 and did not significantly differ from the α-C99/β-C83 ratio found in the nonsense controls. Because a proportionally equal increase of both C-terminal fragments could result either from equally enhanced cleavage at the α- and β-site or reduced γ-cleavage, we also determined Aβ after RNAi down-regulation of calsyntenin-1. Because it has been reported that a substantial proportion of the intracellular Aβ pool is released into the extracellular space...
Fig. 6. Downregulation of calsyntenin-1 by RNAi results in enhanced APP processing and increased secretion of Aβ. Cortical neurons were infected with rAAVs expressing a hairpin targeting calsyntenin-1 (shCst1) or a nonsense hairpin (shNS), together with eGFP. (A) Western blot of cell lysates. The signal indicated equal expression of the recombinant proteins. (B) Quantification of Western blots. Values indicate mean±s.e.m. (*, P<0.05, **, P<0.005). (C–E) Measurement of Aβ secretion. (C) Cortical neurons were infected with rAAV-shCst1 or rAAV-shNS. At DIV12 new medium with DAPT (2 µM) or DMSO carrier was added. Analysis was at DIV14. (D) ELISA of secreted Aβ (1–40) from four experiments. Values indicate mean±s.e.m., n=14 (DMSO) and n=13 (DAPT); **, P<0.005; ***, P<0.001, paired t-test. (E) Immunoblot showing the CTFs of calsyntenin-1 and APP at DIV14. Syntaxin13, Rab11, and β-actin served as loading controls. f, full-length; st, transmembrane stump; β-C99, C-terminal transmembrane stump after γ-secretase cleavage; N=C83, C-terminal transmembrane stump after α-secretase cleavage; N=O-APP695, N- and O-glycosylated mature form of APP695; N-APP695, high-mannose N-glycosylated immature form of APP695; rAAV, recombinant adeno-associated virus containing an expression cassette of the shRNA construct; DAPT, γ-secretase inhibitor DAPT; DMSO, dimethyl sulfoxide.

(Haass and Selkoe, 2007), we quantified Aβ levels in neuronal culture medium after down-regulation of calsyntenin-1, using ELISA (Fig. 6C–E). Compared to control cultures, Aβ secretion was significantly increased by 13% from 260±16 pg/ml to 296±21 pg/ml, P<0.005 (Fig. 6D). To test whether the detected Aβ (1–40) was attributable to release from live neurons we compared release in the absence and presence of the γ-secretase inhibitor DAPT. DAPT reduced the amount of released Aβ by 77%, P<0.001 (Fig. 6D) and resulted in the accumulation of calsyntenin-1 and APP CTFs (Fig. 6E). The increase of Aβ argued against inhibition of γ-secretase as a consequence of calsyntenin-1 down-regulation. Protein levels of ADAM10, BACE1, L1-CAM, syntaxin13, Rab11, and β-actin were identical throughout experimental treatments. We concluded that calsyntenin-1 protects APP from proteolytic processing by its α- and β-secretases during axonal transport.

**Discussion**

Immunoisolated trafficking organelles comprise a combination of multiple vesicular subpopulations – a methodological note

Trafficking organelles are elements of a highly dynamic system of numerous complex and interconnected pathways. The contacts, fusions, and scissions of distinct organelles result in the exchange of “organelle-specific” components. To maintain the proper composition of organelles, components “lost in action” are returned to the location of their function by recycling pathways (Maxfield and McGraw, 2004). This explains why most, if not all of the well characterized “vesicular marker proteins” are also found at lower concentrations in vesicles fusing to or leaving from the respective organelle, as well as in vesicles serving as recycling pathways to maintain the specific molecular inventory of the respective vesicle.

Vesicular cargo proteins are shipped along their intracellular path from one organelle to the other. Therefore, vesicle immunoisolations based on cargo proteins represent a mixture of organelles that are visited by the cargo protein along its trajectory. Also, when vesicles are isolated based on a bona fide “specific” structural or functional component of a trafficking organelle, such as syntaxin13 or Rab5, a clean population of the respective organelle cannot be expected.

Attributable to the composite nature of most immunoisolates, the detection of a given protein by mass spectrometry or Western blotting does not allow for its unequivocal localization. To overcome this drawback, we generated immunoisolates based on partially overlapping vesicular markers. This approach allowed us to more specifically assign the localization of a protein, based on its presence or absence in two or more partially overlapping populations.

The axonal calsytenin-1/APP (Rab11-negative) vesicles represent an anterograde axonal route for delivery of APP and BACE1 to the rapid (early-endosomal) APP recycling pathway of the growth cone

Live-imaging demonstrated that calsytenin-1-containing organelles move in anterograde direction along axons, based on a specific interaction of calsytenin-1 with Kinesin-1 (Konecna et al., 2006; Araki et al., 2007). Subsequent proteomic and immunocytochemical characterizations revealed that the early-endosomal fraction of calsytenin-1 organelles contains APP.
In extension of these studies, we showed here that BACE1 colocalized with calsyntenin-1, APP, and syntaxin13 along axons and in growth cones. We propose that calsyntenin-1, APP, and BACE1 are sorted at the TGN for common post-Golgi anterograde transport. This view is supported by our previous data in COS7 cells, in which calsyntenin-1 and APP intimately colocalized in the TGN and left the TGN together via the formation of tubular structures (Ludwig et al., 2009).

Large pleiomorphic structures in the C-region of growth cones represent the distal end point of the axonal calsyntenin-1/APP trajectory (Steuble et al., 2010). In accordance, live organelle tracking revealed the fusion between calsyntenin-1/APP-tubular structures emerging from the TGN and syntaxin13-labeled structures with early-endosomal morphology (Ludwig et al., 2009). Together, these results strongly suggest that the combined anterograde axonal transportation of APP and BACE1 in calsyntenin-1/Kinesin-1-powered vesicles ends in sorting endosomes of the C-region of the growth cone. Such sorting endosomes have been characterized as a major hub for the peripheral recycling of adhesive and growth-promoting cell surface proteins of the growth cone (Kamiguchi and Lemmon, 2000; Kamiguchi, 2003).

Calsyntenin-1/APP organelles shelter APP during anterograde axonal transport from contact with ADAM10 and protect it from co-transported BACE1. Both organelle immunoisolation and immunocytchemistry characterized the calsyntenin-1/APP vesicle as a carrier in which APP is sheltered from ADAM10. We previously found that calsyntenin-1 is essential for the formation of the calsyntenin-1/APP carrier in the TGN and speculated that the absence of calsyntenin-1 may result in leakage of APP into other organelles. Calsyntenin-1/Kinesin-1-mediated transport from contact with ADAM10 and protect it from co-transported BACE1. Both organelle immunoisolation and immunocytchemistry characterized the calsyntenin-1/APP vesicle as a carrier in which APP is sheltered from ADAM10. We previously found that calsyntenin-1 is essential for the formation of the calsyntenin-1/APP carrier in the TGN and speculated that the absence of calsyntenin-1 may result in leakage of APP into other organelles. Calsyntenin-1/Kinesin-1-mediated transport from contact with ADAM10 and protect it from co-transported BACE1. Both organelle immunoisolation and immunocytchemistry characterized the calsyntenin-1/APP vesicle as a carrier in which APP is sheltered from ADAM10. We previously found that calsyntenin-1 is essential for the formation of the calsyntenin-1/APP carrier in the TGN and speculated that the absence of calsyntenin-1 may result in leakage of APP into other organelles. Calsyntenin-1/Kinesin-1-powered vesicles ends in sorting endosomes of the C-region of the growth cone. Such sorting endosomes have been characterized as a major hub for the peripheral recycling of adhesive and growth-promoting cell surface proteins of the growth cone (Kamiguchi and Lemmon, 2000; Kamiguchi, 2003).

We also found that a substantial proportion of BACE1 destined for EEs of the growth cone is transported along axons together with APP. Our results are in accordance with a study indicating a common Kinesin-mediated transport compartment for BACE and APP in axons of the sciatic nerve, the dorsal root ganglia, and the corpus callosum (Kamal et al., 2001). In contrast, a study in axons of retinal ganglion cells indicated separate transport of YFP-APP and CFP-BACE1 (Goldsbury et al., 2006). Possible explanations for such apparently conflicting observations include differences in the molecular cargo of the distinct trafficking organelles in different neuronal populations, as well as experimentally induced alterations in the molecular cargo due to overexpression of tagged proteins.

The colocalization of BACE1 and APP during calsyntenin-1-mediated axonal transport suggested that β-cleavage of APP may occur before it reaches the growth cone. However, several studies identified the early endosome as the major site of β-cleavage of APP that was imported via endocytosis from the plasma membrane (Koo and Squazzo, 1994; Perez et al., 1999; Kinosita et al., 2003; Rajendran et al., 2006; Rajendran et al., 2008; He et al., 2007). Consistent with this concept, we found predominantly full-length APP in calsyntenin-1, APP, and BACE1 immunoisolates, indicating that APP is not or only slowly processed during axonal transport. This implies the existence of a mechanism that protects APP from proteolysis en route to the growth cone. Several pieces of circumstantial evidence point to calsyntenin-1 as an essential protective agent. Firstly, calsyntenin-1 and APP co-localize from TGN exit to the early endosomal sorting compartment in the C-domain of the growth cone. Secondly, the EE, the major site of APP cleavage by BACE1, coincides with the endpoint of calsyntenin-1/APP colocalization through proteolytic cleavage of calsyntenin-1. Thirdly, knock-down of calsyntenin-1 via RNAi markedly increased the relative amounts of β-CTFs of APP and the secretion of Aβ. Fourthly, APP cleavage in the EE was reported to depend on its ongoing endocytosis from the plasma membrane (Perez et al., 1999; Carey et al., 2005; Cirrito et al., 2008), indicating that endocytosed APP is preferred over anterogradely delivered APP for BACE1 cleavage in EEs.

The mechanism that keeps BACE1 away from APP or that inhibits its activity during its calsyntenin-1/Kinesin-1-mediated anterograde transport is open to conjecture. A mechanism controlling BACE1 activity via redistribution between lipid domains could involve the X11/Mint proteins. The interaction of X11/Mint proteins with APP was found to slow down cellular APP processing (Borg et al., 1998). The formation of a tripartite complex of X11β/Mint2 bound directly to the cytoplasmic segments of calsyntenin-1 and APP was subsequently shown to retard APP processing (Araki et al., 2003).

The reticulon family, in particular reticulon-3 and reticulon-4/Nogo, may directly inactivate BACE1 (He et al., 2004; Murayama et al., 2006). Similarly, complex formation with CSS, the copper chaperone for superoxide dismutase, could inactivate BACE1 (Angeletti et al., 2005). Recent reports indicate an interaction between X11/Mint proteins and CSS (McLoughlin et al., 2001). Based on these observations, it has been suggested that BACE1 could be connected to the APP/X11-complex via CSS and that the formation of a neuronal APP/X11/CSS/BACE1 complex could have an inhibitory effect on BACE1 (Miller et al., 2006). Combining these views, it is tempting to speculate that calsyntenin-1 may interact with the APP/X11/CSS/BACE1 complex in a way that enhances its inhibitory effect on BACE1. Our analyses identified several components of these BACE1-inactivating mechanisms in calsyntenin-1 vesicles. We identified reticulon-4/Nogo by mass spectrometry and X11β/Mint2 by Western blotting (Steuble et al., 2010). Therefore, it is plausible that the BACE1-inhibiting effect of these compartments maintains APP in full-length form during its calsyntenin-1-mediated anterograde axonal trajectory.

Cleavage of calsyntenin-1 by endocytosed ADAM10 may terminate the protection of APP and release axonally transported APP into the peripheral cycling pathway. Our studies which aimed at cell-surface biotinylation of calsyntenin-1 failed despite numerous attempts (not shown), thus suggesting that calsyntenin-1 was proteolytically processed in an internal compartment or within a very short time after its arrival at the plasma membrane. Similarly, the striking clearance of calsyntenin-1 from APP vesicles at the transition from the C- to the P-domain of growth cones suggested an internal compartment as a major site of calsyntenin-1 proteolysis (Steuble et al., 2010). Based on these and the results of our
vesicle incubation studies demonstrating that calsyntenin-1 is cleaved in syntaxin13 immunoisolates, but not in Rab11 and APP immunoisolates, we concluded that calsyntenin-1 is cleaved in early endosomes. Accordingly, most cleaved calsyntenin-1 was found in Rab5 immunoisolates.

A recent report indicated that both ADAM10 and ADAM17, but not BACE1, are capable of cleaving calsyntenin-1 (Hata et al., 2009). Because ADAM17 was reported to be expressed in astrocytes and endothelial cells, but not in neurons (Goddard et al., 2001), we suspected ADAM10 to be the principle terminator of the calsyntenin-1-mediated protection of APP along anterograde axonal transport. We found a relatively strong immunoreactivity for active ADAM10 in the domain of growth cones, thus replenishing full-length APP lost in the rapid recycling endosomal pathway of the growth cone by proteolysis. Calsyntenin-1/Rab11/Rip11 transport packages (red) provide an independent APP-free anterograde transport route, possibly corresponding to the anterograde axonal leg of the long recycling pathway (Lastecka and Winckler, 2011). ADAM10 is transported on a calsyntenin-1-independent transport route. TGN, trans-Golgi network; SV, secretory vesicle; RV, calsyntenin-1-containing early-endosomal replenishment vesicle; RE, calsyntenin-1-containing recycling-endosomal vesicle; EE, early endosome.

Materials and Methods

Antibodies and reagents

Polyclonal rabbit anti-calsyntenin antibodies R85 and R140 have been described earlier (Koncena et al., 2006; Steuble et al., 2010). Anti-Rip11 was provided by Mitsuo Tagaya (Tokyo University of Pharmacy and Life Science) and anti-L1-CAM was provided by Vance Lemmon (University of Miami). Anti-nicastrin (Ab3444) was from Abcam, Cambridge, UK. Anti-GM130 (610822), anti-Mint2 (76120), anti-Rab1b (6100656), anti-Rab4 (610888), anti-syntaxin4 (610439), and anti-syntaxin6 (610635) were from BD Biosciences, Allschwil, Switzerland. Anti-ADAM10 (422751) was from Calbiochem, La¨ufelfingen, Switzerland. Anti-APP (MAB348), anti-PS1 (MAB5232), and anti-Tau-1 (MAB3420) were from Chemicon, Lucerne, Switzerland. Anti-Rab11 (71-5300) was from Invitrogen, Basel, Switzerland. Anti-GFP (1181446001) was from Roche, Basel, Switzerland. Anti-Pen-2 (36-7200), anti-Grp78 (Sc-1050), anti-Mint2 (Sc-30135), anti-PS1 (Sc-7860), anti-Rab5a (Sc-309), and anti-Rab5b (Sc-46692) were from Santa Cruz Biotechnology, Nunningen, Switzerland. Anti-ADAM10 (A2726), anti-APP (A8717) and anti-Tg-actin (A5316) were from Sigma-Aldrich, Buchs, Switzerland. Anti-syntaxin13 (110132), anti-syntaxin16 (110162), anti-VAMP3 (104211), anti-VAMP4 (136002), and anti-VSVG (169022) were from Synaptic Systems, Göttingen, Germany. Anti-BACE1 (PA1-757) was from Thermo Scientific, Wohlen, Switzerland. Fluorescent secondary antibodies (Cy3-, FITC-, Cy5-, and DyLight649-conjugated) were from Jackson ImmunoResearch Laboratories, West Grove, PA, USA and used at 2.5 μg/ml.

The γ-secretase inhibitor DAPT (1 mM in DMSO) and TPEN, a cell-permeable inhibitor of Zn2+-dependent matrix metalloproteases (10 mM in DMSO) were purchased from Sigma-Aldrich. The β-secretase inhibitor IV (20 mM in DMSO) was purchased from Calbiochem. N-glycosidase F and endoglycosidase H were from Roche.

Subcellular fractionation of mouse brain and immunoinositolization of vesicular organelles

The V1 membrane fraction was prepared from P7 mouse brains by differential centrifugation as previously described (Morfini et al., 2002; Koncena et al., 2006; Steuble et al., 2010). Washed V1 pellets were resuspended in IP buffer (PBS, 320 mM sucrose, 5 mM EDTA, pH 7.4) and stirred for 1 h at 4°C. For immunoblotting, 2 mg magnetic Dynabeads M-280 protein A (Invitrogen) were preincubated with 10 μg IgG for 40 min and washed four times in IP buffer. V1 inputs were adjusted to ~0.7 mg/ml with IP buffer and incubated with pre-coated beads for 1 h at 4°C. Beads with immunoinosolated organelles were washed 10 times with 1 ml IP buffer, once with 20 mM Tris·Cl, pH 7.4, and subsequently incubated in 50 μl 20 mM Tris·Cl, pH 7.4, 0.1% (v/v) Triton X-100 for 30 min at 25°C.

For Western blotting, 20 μg protein from input and 40 μl eluate were resolved on 4–12% NuPage Bis-Tris gels (Invitrogen). Bead contents were analyzed separately. Immunoblots were imaged with a Fuji LAS-3000 Lite CCD camera (Raytest AG, Wetzikon, Switzerland) and quantified using the AIDA 2D multi labeling software (version 3.4; Raytest AG).
For in vitro incubation experiments, immunosolated organelles were incubated for 30 min at 37°C in PBS, pH 7.4, 2 mM MgCl2, 3 mM ATP, supplemented with vehicle or 2 μM DAPT, 20 μM TPEF, and 10 μM β-secretase inhibitor IV, respectively (Yamashiro et al., 1983).

Cultures of dissociated hippocampal and cortical neurons

For immunocytochemistry, 5,000 dissociated hippocampal cells/cm2 from E19 NMRI mice were plated onto poly-L-lysine-coated (Sigma-Aldrich) glass coverslips and co-cultured face-to-face with a monolayer of astrocytes (Banker, 1980). For Western blotting, 500,000 dissociated cortical cells/cm2 were plated onto poly-L-lysine-coated 12-well plates. Cells were grown in neurobasal medium supplemented with B27, 5 mM glutamine, and antibiotics (penicillin-streptomycin).

Immunocytochemistry

For immunofluorescence analysis, neurons were fixed in 4% paraformaldehyde, 4% sucrose in PBS, pH 7.4, for 10 min at room temperature. Samples were blocked for 1 h in 10% fetal calf serum, 0.1% saponin in PBS, pH 7.4. Neurons were then exposed to primary antibodies in 3% fetal calf serum, 0.1% saponin in PBS, pH 7.4, overnight at 4°C, then incubated for 1 h with Cy3-, FITC-, Cy5-, or DyLight649-conjugated secondary antibodies and mounted in Vectashield medium (Vector Laboratories, Burlingame, CA, USA).

In some experiments, double-labeling had to be carried out with two primary antibodies raised in the same species. In this case, neurons were first labeled with the first primary antibody, washed, and incubated with an excess of goat anti-rabbit Fab fragments (1:10) following the manufacturer’s instructions (Jackson ImmunoResearch Laboratories). After extensive washing, cells were incubated with the second primary antibody, followed by incubation with anti-goat and anti-rabbit secondary antibodies. Two control experiments confirmed the specificity of this approach. Firstly, it was shown that FITC-conjugated anti-rabbit secondary antibody was not immunoreactive in the presence of primary rabbit antibody masked with goat Fab fragments. This control served to stringently define the confocal settings during imaging and image processing. Secondly, the staining patterns of double-labeled cells were shown to be identical to immunostainings of the respective antibodies alone.

Confocal images were taken with a Leica confocal laser scanning microscope TCS-SP2 at a resolution of 1024×1024 pixels using a Leica PL Apo 63× (NA=1.32) objective (Leica, Heerbrugg, Switzerland). Colocalization was analyzed by ImageJ (version 1.38x; National Institutes of Health, Bethesda, MA, USA). Statistical analyses were performed with Prism (version 4.0; GraphPad Software Inc, La Jolla, CA, USA). RGB images were used to determine the degree of colocalization between the red, green, and blue channels using the colocalization RGB plugin of ImageJ. Puncta were counted from binarized images with the particle analyzer tool, minimum size of counted puncta set to 20 pixels. Data shown are mean±s.e.m. of at least four separate images.

Generation of recombinant mouse adenovirus-associated viruses (AAV) for RNAi

To generate calnsyn-1 small hairpin knockdown construct pBlueU6-siCst1-1, the promoter of the mouse U6 RNA gene (GenBank X06980) was amplified from genomic DNA using oligonucleotides GCGGATCCGACGCCATCTCTA and GCTCTAGAGGAGTTAAACAGGCTTTTCTCCAAG. The resulting PCR product was digested with BamHI and XbaI and cloned into pBluescript II SK (+) (Stratagene). A 19 nt sequence targeting mouse calsyntenin-1 (548–476 bp; GCTCTAGAGCGTTAACAAGGCTTTTCTCCAAGG. The resulting PCR product was amplified by using the respective antibodies alone.

Generation of recombinant adenovirus

APP-mRFP, BACE-eGFP, meGFP-Cst-1, and mRFP-Cst-1 were subcloned into a transfer vector to generate infectious adenovirus H5, as previously described (Frischknecht et al., 2008). Viral particles were purified by a single centrifugation over a CsCl step gradient (1.25 and 1.4 g/ml CsCl). Expression of recombinant proteins was assessed by immunoblotting of lysates of infected HEK-293 cells.

Live-imaging of axonal transport of fluorescently-tagged calsynin-1, APP, and BACE1

Cultures of dissociated hippocampal neurons were co-infected with adenovirus on DIV7 and imaged after two days in a Ludin imaging chamber (Life Imaging Services, Basel, Switzerland) under constant perfusion with (in mM): 119 NaCl, 2.5 KCl, 2 CaCl2, 2 MgCl2, 30 glucose, and 25 HEPES, pH 7.4. Widefield images were acquired on a Leica LX microscope equipped with a 100×, 1.4 NA, oil objective, a Hamamatsu-C9100-13 EM-CCD camera system (500×500 px, Hamamatsu, Horsching am Ammersee, Germany). Images were taken at 1 sec intervals for a period of 1 min. Axons were identified according to morphological criteria (thin and at least two times longer than dendrites). Anterograde and retrograde vesicle transport was quantified independently for each channel and subsequently tested for colocalization. The quantification was performed manually and included only vesicles that were clearly traceable for at least 20 sec.

ELISA for murine Aβ

Cortical neurons were infected with the respective rAAVs on the day of plating and kept with virus for 12 DIV. Neurobasal medium was removed and neurons were washed twice with PBS, pH 7.4, to remove Aβ that had accumulated during cultivation. Then, 300 µl of fresh medium without phenol red, supplemented with B27, vehicle or 1 μM DAPT were added. After 2 days, 250 µl medium were harvested and centrifuged for 30 min at 100,000×g at 4°C. Samples (100 µl) were added to a 96-well ELISA plate (code 27720; Immuno-Biological Laboratories, Minneapolis, MN, USA) and processed according to the manufacturer’s instructions.

Glycosidase treatments in vitro

V1 membranes or syntaxin13 immunosolates were processed with N-glycosidase F (PNGaseF, Roche) or endoglycosidase H (EndoH, Roche). For deglycosylation with PNGaseF, the reaction mixture (80 µl final volume) contained 10 µg protein and 4 µl PNGaseF in 10 mM EDTA, 1% β-mercaptoethanol (v/v), 0.1% SDS, and 2.75 mM Tris-Cl, pH 7.4. For deglycosylation with EndoH, the reaction mixture (80 µl final volume) contained equal amounts of protein and 2 µl EndoH in 1% β-mercaptoethanol (v/v), 0.25% SDS, and 10 mM sodium acetate, pH 5.2. Enzymes were added for 16 h at 37°C after denaturation of protein samples at 95°C for 5 min.

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Competing interests

The authors declare that there are no competing interests.

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