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Calsyntenin-1 mediates axonal transport of the amyloid precursor protein and regulates A β production

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Understanding the mechanisms that control processing of the amyloid precursor protein (APP) to produce amyloid- β (A β) peptide represents a key area of Alzheimer's disease research. Here, we show that siRNA-mediated loss of calsyntenin-1 in cultured neurons alters APP processing to increase production of A β . We also show that calsyntenin-1 is reduced in Alzheimer's disease brains and that the extent of this reduction correlates with increased A β levels. Calsyntenin-1 is a ligand for kinesin-1 light chains and APP is transported through axons on kinesin-1 molecular motors. Defects in axonal transport are an early pathological feature in Alzheimer's disease and defective APP transport is known to increase A β production. We show that calsyntenin-1 and APP are co-transported through axons and that siRNA-induced loss of calsyntenin-1 markedly disrupts axonal transport of APP. Thus, perturbation to axonal transport of APP on calsyntenin-1 containing carriers induces alterations to APP processing that increase production of A β . Together, our findings suggest that disruption of calsyntenin-1-associated axonal transport of APP is a pathogenic mechanism in Alzheimer's disease.

INTRODUCTION

Deposition of amyloid- β (A β) within amyloid plaques is a hallmark pathology of Alzheimer's disease. A β is an approximate 40 amino acid peptide that is derived by proteolytic cleavage from amyloid precursor protein (APP). Processing of APP to produce A β involves cleavage by β -site APP cleaving enzyme-1 (BACE1) and γ -secretase that process APP at the N- and C-termini, respectively, of the A β sequence. In addition, APP can be cleaved by α - and γ -secretases and this precludes A β production since α -secretase cleaves APP within the A β sequence (1).

A large body of evidence suggests that altered production of A β is a major pathogenic event in Alzheimer's disease (2). Indeed, some familial forms of Alzheimer's disease are caused by mutations in the *APP* gene and several of these

mutations alter processing of APP and production of A β (2). Understanding the molecular mechanisms that control APP processing thus represents a key area of Alzheimer's disease research. Alterations to APP trafficking are acknowledged to be one mechanism for modulating APP processing and A β production (3).

Neurons are especially dependent upon correct protein and organelle trafficking since they are polarized with axons and dendrites, and also because transport through axons can involve cargo movement over exceptionally long distances. Moreover, a large body of evidence now implicates defective axonal transport in Alzheimer's disease (reviewed in 4–6). Within neurons, APP is synthesized in cell bodies and then undergoes anterograde axonal transport on kinesin-1 molecular motors (7,8). Most functional kinesin-1 comprises a heterotetramer of two kinesin-1 motor proteins and two kinesin-1

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light chains (KLCs). Kinesin-1 contains ATPase activity and uses the chemical energy of ATP to drive conformational changes that generate motile force; in contrast, the KLCs are mainly involved in binding of cargoes (9).

The precise mechanisms by which APP attaches to and is transported by kinesin-1 are not properly understood (reviewed in 10). However, there is evidence that one route may involve calyntenin-1 (also known as alcadin- α) (11,12). Calyntenin-1 is a neuronally enriched type-1 membrane-spanning protein that binds directly to KLCs via its intracellular C-terminal domain (13–16). As such, calyntenin-1 acts as a ligand to mediate transport of a subset of vesicles through axons on kinesin-1 motors. A proportion of APP and calyntenin-1 co-localize in cells and tissues, and proteomic studies have shown that some calyntenin-1 containing vesicles also contain APP (11,12). However, such studies are correlational and do not formally demonstrate that calyntenin-1 is required for movement of APP in neurons. Indeed, other studies suggest that calyntenin-1 does not normally mediate axonal transport of APP (14). Likewise, the effect of calyntenin-1 on APP processing and A β production is unclear. Some studies indicate that the loss of calyntenin-1 promotes APP processing, whereas others indicate that over-expression of calyntenin-1 increases A β production (12,14). Notably, the effect of loss of calyntenin-1 on A β production in neurons has not been reported.

To obtain formal evidence on the role of calyntenin-1 in transport and processing of APP, we monitored how siRNA loss of calyntenin-1 influenced APP axonal transport and production of endogenous A β in living neurons. Here, we show that APP and calyntenin-1 are co-transported through axons, that the loss of calyntenin-1 disrupts axonal transport of APP and that calyntenin-1 loss also leads to altered APP processing and increased production of A β . Finally, we demonstrate that calyntenin-1 expression is reduced and negatively correlates with A β burden in Alzheimer's disease brains. Thus, altered axonal transport of APP on calyntenin-1 carriers may be mechanistic in Alzheimer's disease.

RESULTS

siRNA-mediated loss of calyntenin-1 increases A β production in neurons

We first tested the role of calyntenin-1 on production of A β . To do so, we downregulated calyntenin-1 expression in rat cortical neurons using siRNAs and monitored production of endogenous A β (1–40) and A β (1–42) species using ELISAs. Two different siRNAs and a Smartpool mix of four siRNAs all reduced calyntenin-1 expression to lower than 5% of that seen in control cells (Fig. 1A). A β ELISAs revealed that this reduction in calyntenin-1 expression led to a significant approximate 1.65-fold increase in secretion of both A β (1–40) and A β (1–42) (Fig. 1B). Since all three siRNAs produced highly similar effects on calyntenin-1 expression and A β production, the Smartpool mix was used in all later experiments.

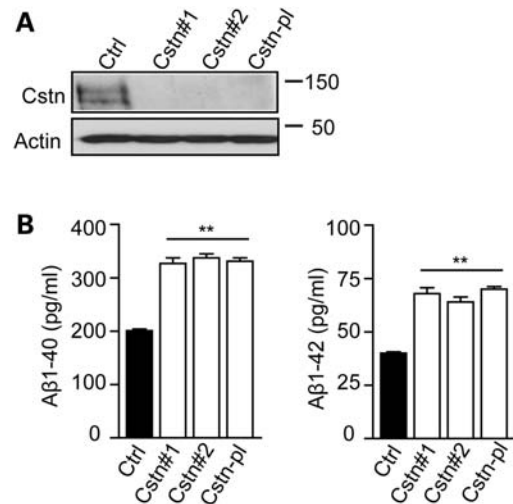


Figure 1. siRNA knockdown of calyntenin-1 increases production of A β . (A) Knockdown of calyntenin-1 in rat cortical neurons. Neurons were treated with control siRNA (Ctrl) or with two different calyntenin-1 siRNAs (Cstn#1; Cstn#2) or with a pool of four different calyntenin-1 siRNAs (Cstn-pl). Samples were then probed on immunoblots for calyntenin-1 (Cstn) or actin as a loading control. Molecular mass markers are shown. (B) ELISA assays for A β (1–40) and A β (1–42) in conditioned media from rat cortical neurons treated with control siRNA (Ctrl) or with two different calyntenin-1 siRNAs (Cstn#1; Cstn#2) or with a pool of four different calyntenin-1 siRNAs (Cstn-pl). Treatment with calyntenin-1 siRNAs increase the amounts of both A β (1–40) and A β (1–42) in conditioned media. Statistical significance was determined by one-way ANOVA followed by the Bonferroni *post hoc* test. $n = 6$; error bars are SEM; ** $P < 0.01$.

siRNA-mediated loss of calyntenin-1 increases APP processing at the BACE1 site but does not alter the levels of BACE1, ADAM10, presenilin-1, nicastrin, full-length APP or phosphorylation of APP on threonine-668

To gain insight into the mechanisms by which the loss of calyntenin-1 increased production of A β , we first monitored APP levels and APP processing at the α -secretase and BACE1 sites in neurons treated with control or calyntenin-1 siRNAs by immunoblotting. Processing of APP by α -secretase and BACE1 generates N-terminal secreted APP fragments (sAPP α and sAPP β) and the corresponding membrane-associated APP C-terminal domains. These APP C-terminal domains correspond to the two fragments produced by BACE1 processing (99 and 89 amino acids; APP-C99 and APP-C89) and the fragment produced by α -secretase processing (83 amino acids; APP-C83) (1). While the C-terminal APP fragments can be resolved on SDS-PAGE, analysis is complicated since in neurons, a proportion of APP is phosphorylated on threonine-668 (APP-thr668) by c-Jun N-terminal kinase-3 which alters the migration pattern of the different processed species so that some co-migrate (17,18). However, treatment of samples with λ -protein phosphatase prior to analyses facilitates interpretation of the band pattern (18).

Loss of calyntenin-1 did not alter the total levels of full-length APP (Fig. 2A). However, analyses of APP C-terminal processed fragments (APP-CTFs) in cell lysate samples showed that the loss of calyntenin-1 decreased the level of APP-C83 and increased the levels of both APP-C89 and

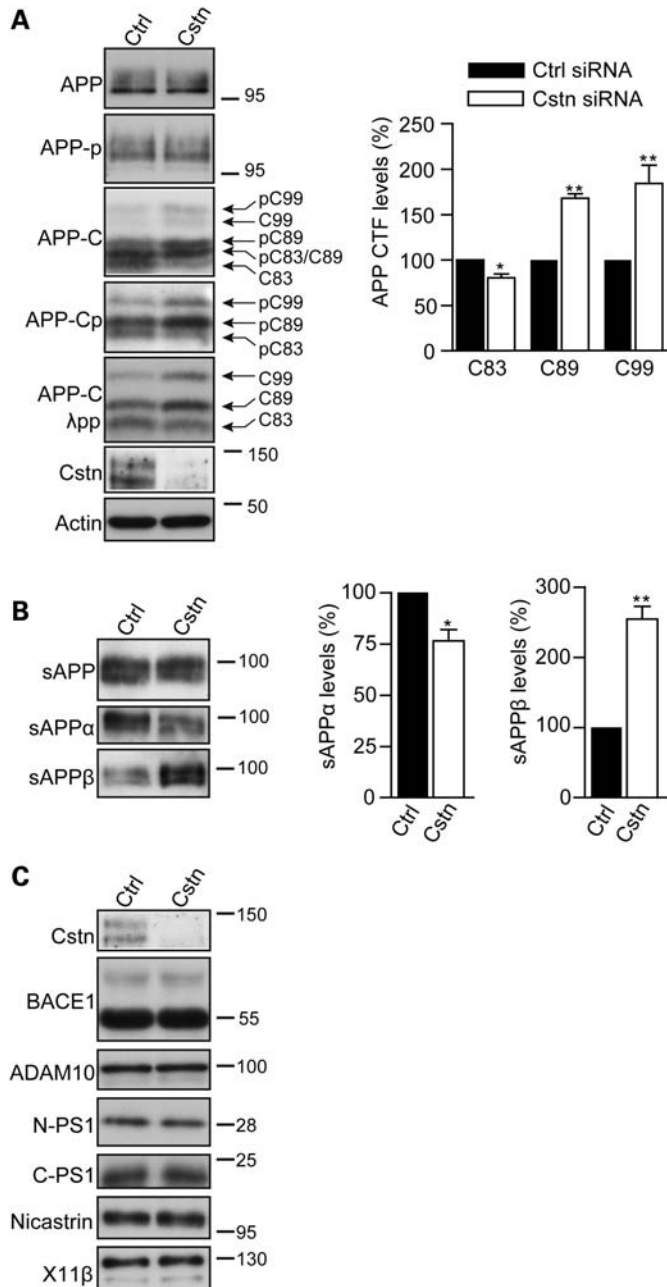


Figure 2. siRNA knockdown of calyntenin-1 alters APP processing in rat cortical neurons without affecting expression of APP or APP secretase proteins. (A) Neurons were treated with control (Ctrl) or calyntenin-1 (Cstn) siRNAs (pooled mix of four different siRNAs) and samples then probed on immunoblots for full-length APP (APP), full-length APP phosphorylated on threonine-668 (APP-p), APP C-terminal processed fragments (APP-C), APP C-terminal processed fragments phosphorylated on threonine-668 (APP-Cp) or APP C-terminal processed fragments following treatment with λ -protein phosphatase (APP-C λ pp). Immunoblots showing APP C-terminal processed fragments are marked to indicate APP species generated following cleavage by BACE1 (C99 and C89) or α -secretase (C83), and their phosphorylated variants (pC99, pC89, pC83). Bar graphs show relative levels of APP C-terminal fragments (APP-C83; APP-C89 and APP-C99) in the different samples following treatment with λ -protein phosphatase. Statistical significance was determined by Student's *t*-test. $n = 5$; error bars are SEM. * $P < 0.05$; ** $P < 0.01$. No significant changes in the levels of full-length APP or full-length APP phosphorylated on threonine-668 were detected in the same samples. (B) sAPP fragments in conditioned media of rat cortical neurons

APP-C99 fragments (Fig. 2A). These changes were most easily observed in the λ -protein phosphatase-treated samples (Fig. 2A). To complement these analyses, we also monitored APP processing by immunoblotting for APP-secreted fragments in conditioned media. Probing with antibody 22C11 (that detects total sAPP) and antibodies that specifically detect sAPP α and sAPP β revealed that while the loss of calyntenin-1 did not alter the total sAPP levels, it induced a decrease in sAPP α and a corresponding increase in sAPP β levels (Fig. 2B).

APP-thr668 phosphorylation has been linked to axonal transport of APP and A β production (19–21). We therefore monitored the effect of calyntenin-1 knockdown on APP-thr668 phosphorylation. Loss of calyntenin-1 did not alter the levels of phosphorylated full-length APP (Fig. 2A). Moreover, changes in phosphorylation of APP-C99, APP-C89 and APP-C83 were in line with the relative changes in the total amounts of these processed fragments (Fig. 2A).

We also monitored whether the loss of calyntenin-1 altered the levels of BACE1, the α -secretase ADAM10, the γ -secretase components presenilin-1 and nicastrin (1) and X11 β by immunoblotting. X11 β (also known as X11-like or munc-18 interacting protein-2) binds to both APP and calyntenin-1 (22,23). Knockdown of calyntenin-1 did not change the levels of any of these secretases or X11 β (Fig. 2C). Thus, the increased production of A β associated with the loss of calyntenin-1 is accompanied by a decrease in processing of APP at the α -secretase site and a corresponding increase in processing at the BACE1 sites. However, these changes do not involve alterations to the levels of full-length APP, APP-thr668 phosphorylation, APP secretases or X11 β .

A significant proportion of APP and calyntenin-1 co-localize in axons and APP-mCherry and enhanced green fluorescent protein-calyntenin-1 are co-transported through living axons

Since calyntenin-1 functions as a ligand for kinesin-1 and APP is transported on kinesin-1 motors, the effect of calyntenin-1 loss on APP processing and A β production may be due to changes in axonal transport of APP. Indeed, disruption of kinesin-1-mediated APP axonal transport alters APP processing to increase A β production (8). To begin to test this possibility, we first monitored co-localization of APP and calyntenin-1 in neurons. Both APP and calyntenin-1 are enriched in the Golgi and calyntenin-1 has been shown to mediate kinesin-1 transport of cargoes on post-Golgi carriers (12,15). In line with these findings, a proportion of calyntenin-1 co-localized with the trans-Golgi network (TGN) marker TGN-38 and proportions of both calyntenin-1

treated with control (Ctrl) or calyntenin-1 (Cstn) siRNAs. Samples were probed on immunoblots for total sAPP (sAPP), sAPP α and sAPP β . Bar graph shows relative levels of sAPP α and sAPP β in the different samples; total sAPP levels were unchanged. Statistical significance was determined by Student's *t*-test. $n = 4$; error bars are SEM. * $P < 0.05$; ** $P < 0.01$. (C) siRNA knockdown of calyntenin-1 does not affect expression of BACE1, α - or γ -secretase proteins or X11 β . Neurons were treated with control (Ctrl) or calyntenin-1 siRNAs (Cstn) and samples ($n = 4$) then probed on immunoblots for BACE1, ADAM10, N- and C-terminal presenilin-1 (PS1) fragments, nicastrin and X11 β as indicated. Molecular mass markers are shown.

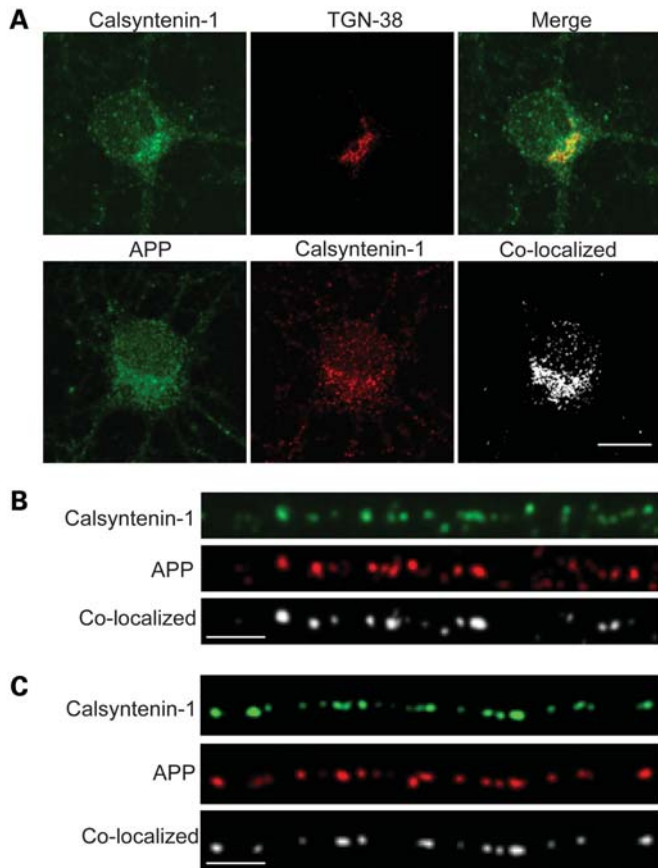


Figure 3. Calyntenin-1 is present in the TGN and APP and calyntenin-1 co-localize in cell bodies and in axons of cultured rat cortical neurons. (A and B) Neurons were co-stained for calyntenin-1 and either TGN-38 or APP as indicated. (A) Confocal slices through cell bodies with merge (calyntenin-1 and TGN-38) or co-localized pixels (calyntenin-1 and APP). (B) Endogenous calyntenin-1 and APP in axons with co-localized puncta. (C) EGFP-calyntenin-1 and APP-mCherry in axons with co-localized puncta. Scale bars are 10 μm (A) and 5 μm (B and C).

and APP co-localized in neuronal cell bodies (Fig. 3A). Some APP and calyntenin-1 also co-localized in vesicular structures in axons (Fig. 3B). Quantification of this axonal co-localization revealed that 41% of APP vesicles co-localized with calyntenin-1. These findings are in broad agreement with previous studies which showed that in sciatic nerve and in axons of cultured hippocampal neurons, ~ 30 –45% of APP and calyntenin-1 co-localize (11,14). We also monitored the axonal distributions of transfected enhanced green fluorescent protein (EGFP)-calyntenin-1 and APP-mCherry in neurons. High levels of co-localization of these transfected proteins were detected in axons (Fig. 3C).

The co-localization of APP and calyntenin-1 in axons suggest that they are co-transported. We therefore monitored axonal transport of APP-mCherry and EGFP-calyntenin-1 by dual imaging time-lapse microscopy in co-transfected living rat cortical neurons. In line with our previous work (15,24), we chose cells expressing low levels of transfected proteins (as judged by the fluorescent protein signal) for analyses so as to avoid any possible artefacts produced by high levels of expression. Co-ordinate movement of APP-mCherry and EGFP-calyntenin-1 in both anterograde and retrograde

directions was observed through axons (Fig. 4A). Movement was predominantly anterograde and mean speeds were 1.6 $\mu\text{m/s}$ (anterograde) and 1.1 $\mu\text{m/s}$ (retrograde). These velocities and the bias towards anterograde movement are similar to those described previously for calyntenin-1 (13–15). APP-YFP/EGFP also moves in a predominantly anterograde direction with reported speeds of 0.8–9 $\mu\text{m/s}$ which may depend to some extent upon the cell type utilized (7,14,25–27). Notably, $\sim 65\%$ of APP-EGFP is transported at speed of 1–2 $\mu\text{m/s}$ in mouse cortical neurons which are similar to the speeds reported here in rat cortical neurons (26).

siRNA loss of calyntenin-1 decreases axonal transport of APP

The above co-localization and live imaging studies suggest that a significant proportion of APP is co-transported with calyntenin-1 through axons. To test further this possibility, we quantified APP-EGFP axonal transport using time-lapse microscopy in transfected living rat cortical neurons following siRNA knockdown of calyntenin-1. APP-EGFP movement was again predominantly anterograde in these neurons. However, analyses of the flux rates for APP-EGFP transport revealed that the loss of calyntenin-1 markedly reduced the numbers of APP-EGFP moving cargoes (Fig. 4B–D). This reduction was seen in both anterograde (64.3% reduction) and retrograde (46.6% reduction) directions (Fig. 4D). Thus, APP and calyntenin-1 co-localize in axons, APP-mCherry and EGFP-calyntenin-1 are co-transported through axons, and the loss of calyntenin-1 markedly disrupts axonal transport of APP.

siRNA loss of calyntenin-1 increases APP levels in the TGN

Calyntenin-1 is enriched in the Golgi and mediates trans-Golgi exit of APP (12). The reduced axonal transport of APP-EGFP seen in calyntenin-1 knockdown neurons suggests that APP may be retained in the cell body and accumulate in the TGN. To test this possibility, we quantified the APP fluorescent signals in the TGN in control and calyntenin-1 siRNA-treated neurons. Loss of calyntenin-1 induced a significant 54.7% increase in APP signals in the TGN (Fig. 5).

Calyntenin-1 expression is reduced in Alzheimer's disease brains and the extent of this reduction correlates inversely with A β levels

The above studies, which show that the loss of calyntenin-1 increases A β production and that this is associated with a disruption to axonal transport of APP, suggest that defects in calyntenin-1 metabolism may be mechanistic in Alzheimer's disease. To begin to test this possibility, we monitored calyntenin-1 expression in Alzheimer's disease brains. Immunoblotting of cortical samples from control and Braak stage V–VI Alzheimer's disease brains revealed that compared with controls, calyntenin-1 expression was significantly reduced in Alzheimer's disease brains ($73.5 \pm 6.72\%$ of control) (Fig. 6A). There was no significant correlation between calyntenin-1 levels and post-mortem delay.

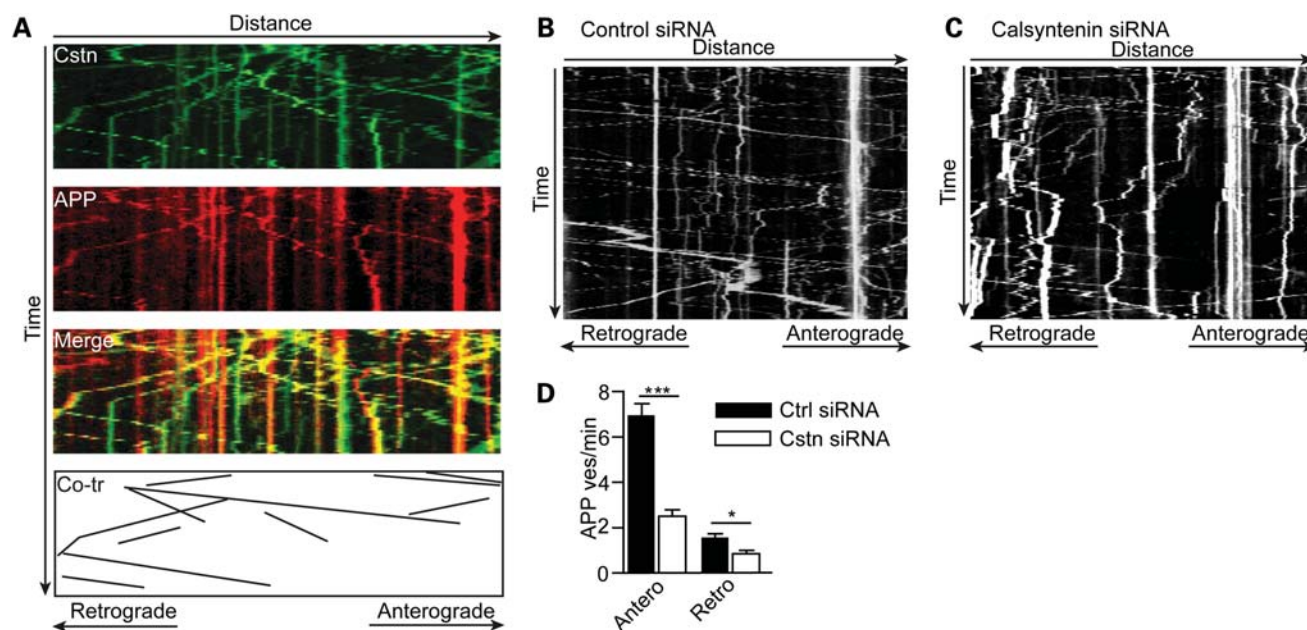


Figure 4. EGFP-calsyntenin-1 and APP-mCherry are co-transported through axons of rat cortical neurons and siRNA loss of calsyntenin-1 disrupts axonal transport of APP-EGFP. (A) Kymograph showing co-transport of EGFP-calsyntenin-1 (Cstn) and APP-mCherry (APP). Merge and co-transported (Co-tr) EGFP-calsyntenin-1/APP-mCherry particles are shown. (B and C) Representative kymographs showing axonal transport of APP-EGFP in control (B) and calsyntenin-1 siRNA-treated neurons. (D) Flux rates for APP-EGFP vesicle movement in control and calsyntenin-1 siRNA-treated neurons. Loss of calsyntenin-1 disrupts APP-EGFP transport in both anterograde and retrograde directions. Statistical significance was determined by Student's *t*-test. $n = 33$ control and 33 calsyntenin-1 siRNA-treated cells; error bars are SEM; * $P < 0.05$; *** $P < 0.001$.

We also tested whether calsyntenin-1 levels correlated with A β burden in these samples. Total A β (1–40) and A β (1–42) were quantified by ELISAs and correlated with calsyntenin-1 protein levels. Significant negative correlations were observed between both A β (1–40) ($r^2 = -0.396$) and A β (1–42) ($r^2 = -0.441$), and calsyntenin-1 levels (Fig. 6B).

DISCUSSION

Here, we show that the loss of calsyntenin-1 increases production of endogenous A β (1–40) and A β (1–42) in rat neurons. We also show that calsyntenin-1 levels are reduced significantly in Alzheimer's disease brains and that the extent of this reduction correlates significantly with increased A β levels. Loss of calsyntenin-1 in the cultured neurons was associated with alterations to APP processing involving increased cleavage at the BACE1 sites and decreased cleavage at the α -secretase site.

APP is synthesized in the endoplasmic reticulum (ER) and then trafficked through the Golgi to the cell surface. Here, it can be re-internalized to endocytic and recycling compartments and trafficked back to the cell surface or degraded in the lysosome (3). Calsyntenin-1 is enriched in the TGN and recent evidence suggests that it mediates transport of a subset of cargoes on post-Golgi carriers via kinesin-1 (11,12). Our findings that siRNA loss of calsyntenin-1 reduces axonal transport of APP and increases APP levels in the TGN are thus consistent with a role for calsyntenin-1 in mediating transport of APP on post-Golgi carriers.

The precise intracellular location(s) where APP is processed to produce A β in neurons are not clear. BACE1 localizes

mainly to the TGN and endosomes, whereas γ -secretase components are found in multiple locations, including ER, Golgi, endosomes and the cell surface (3,28). Thus, some studies suggest that A β is produced in endosomes (29–33), whereas others suggest that the TGN is a primary site for A β production (34–39). There is also evidence that A β is produced in other locations, including autophagosomes (40). Whatever the scenario, disrupting anterograde transport of APP may well promote amyloidogenic processing by targeting APP to sites of cleavage by BACE1 and/or γ -secretase. In particular, our finding that siRNA loss of calsyntenin-1 increases APP levels in the TGN, a favoured site for APP processing by BACE1 (see above) may explain the increased cleavage of APP at the BACE1 sites that we detect in the calsyntenin-1 knockdown neurons.

Despite highly efficient siRNA knockdown of calsyntenin-1, its loss reduced but did not abolish anterograde axonal transport of APP. Other mechanisms for transport of APP by kinesin-1 have been described, including direct attachment of APP to KLCs, or indirect attachment to KLCs via adaptor proteins such as JIPs (C-Jun N-terminal kinase interacting proteins) and PAT1 which bind to both KLC and APP (14,19,41–43). The GTPase Rab3A has also been implicated in APP transport (26). Transport involving these other mechanisms may account for the remaining APP movement seen in the calsyntenin-1 knockdown neurons. It is also possible that the co-ordinate transport of APP-mCherry and EGFP-calsyntenin-1 seen in our dual imaging studies overestimates the proportion of APP that is co-transported with calsyntenin-1 (e.g. overexpression of calsyntenin-1 in these experiments may recruit APP from JIP1/PAT1 carriers).

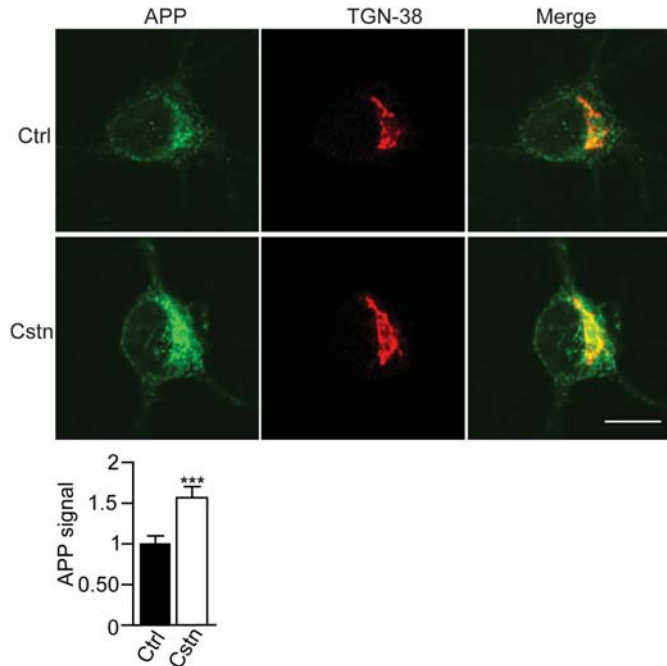


Figure 5. siRNA loss of calyntenin-1 increases APP levels in the TGN. Neurons were treated with control (Ctrl) or calyntenin-1 (Cstn) siRNAs and immunostained for APP and TGN-38. Representative cells are shown with merge. Scale bar is 10 μ m. Bar chart shows relative APP fluorescent signals in the TGN. Statistical significance was determined by Student's *t*-test. $n = 50$ control and 54 calyntenin-1 siRNA-treated cells; error bars are SEM; *** $P < 0.001$.

Nevertheless, our observations that siRNA knockdown of calyntenin-1 markedly inhibits anterograde movement of APP strongly suggests that a significant proportion of APP is transported on calyntenin-1 carriers. Indeed, using alternative approaches that do not directly assay movement of APP through axons, others have also concluded that an important route for axonal transport of APP by kinesin-1 may involve calyntenin-1 (11,12).

Our findings that the loss of calyntenin-1 disrupts axonal transport of APP and increases production of A β focus attention on the mechanism that controls APP transport on calyntenin-1 carriers. In particular, the mechanisms that control loading of APP onto calyntenin-1 containing vesicles and binding of calyntenin-1 to KLCs represent key regulatory points that could influence APP trafficking.

Although the route(s) by which APP is loaded onto calyntenin-1-containing vesicles are not clear, both APP and calyntenin-1 bind to X11 β (22,23,44). X11 β is a neuronal adaptor protein that is enriched in the Golgi and which regulates APP processing and A β production (45–50). There is evidence that X11 β functions as a vesicle coat protein; coat proteins function in the selection of cargo for inclusion into particular vesicle sub-types (51). Interestingly, phosphorylation of KLC1 on serine-460 has recently been shown to selectively regulate binding of calyntenin-1 but not several other ligands to kinesin-1 motors (15). Thus, via its binding to both APP and calyntenin-1, X11 β may function to load APP onto calyntenin-1-containing vesicles for post-Golgi transport on kinesin-1 motors, and phosphorylation of KLCs

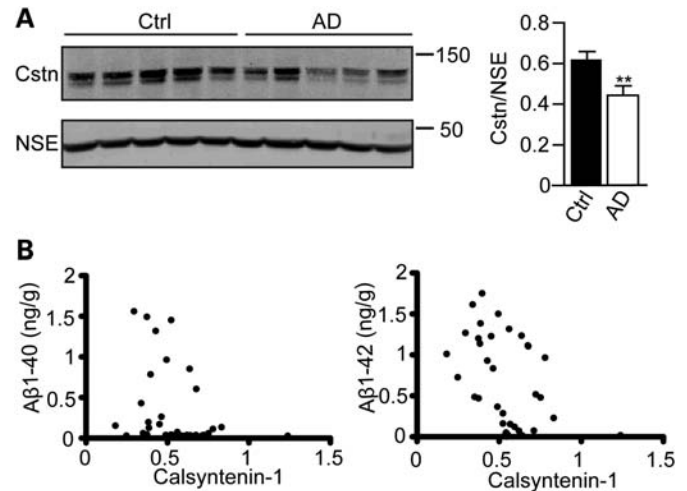


Figure 6. Calyntenin-1 is reduced in Alzheimer's disease brains and the extent of this reduction correlates with A β levels. (A) Immunoblots of representative samples from control (Ctrl) and Alzheimer's disease (AD) cortex. Samples were probed for calyntenin-1 (Cstn) and NSE. Bar graph shows relative levels of calyntenin-1 in the two sample sets following normalization to NSE signals in each sample. Statistical significance was determined by Student's *t*-test. $n = 19$ control (Ctrl) and $n = 16$ Alzheimer's disease samples; error bars are SEM. ** $P < 0.01$. (B) Correlation between cortical calyntenin-1 amounts and A β burden in post-mortem brains. A β levels in human tissues were correlated with calyntenin-1 signals by generating correlation coefficients and significance established by non-parametric, two-tailed, Spearman Rho tests. Graphs show A β (1–40) and A β (1–42) levels (ng/g brain tissue) plotted against calyntenin-1 signals (arbitrary units) obtained from immunoblots. Significant negative correlations were observed between both A β (1–40) ($r^2 = -0.396$; $P < 0.05$) and A β (1–42) ($r^2 = -0.441$; $P < 0.01$) and calyntenin-1 levels ($n = 19$ control; $n = 16$ Alzheimer's disease samples).

may regulate attachment of calyntenin-1 with associated APP to kinesin-1. Disruption to loading of APP onto calyntenin-1 vesicles or transport of APP by kinesin-1 on calyntenin-1 containing vesicles are thus likely to promote amyloidogenic processing of APP. Moreover, the negative correlation we detect between calyntenin-1 and A β levels in Alzheimer's disease brains suggest that the loss of calyntenin-1 is mechanistic in the disease process.

MATERIALS AND METHODS

Plasmids and siRNAs

EGFP-calyntenin-1 and APP-EGFP were as described previously (13,52). APP-mCherry was generated by replacement with mCherry sequences in APP-EGFP. Calyntenin-1 and negative control siRNAs were from Dharmacon (Accell range). Calyntenin-1 siRNAs were #1 5'-GCAAAGAGC AUCAGUAUAA-3', #2 5'-CCCUAAAGAUGUGUAUUAU U-3', #3 5'-CCAUCACGCUUGCAGUUUU-3' and #4 5'-GUGUCAGCUUCCUGUUUU-3'. Negative control siRNA (catalogue number D-001910-03) has been designed, modified and microarray-confirmed by Dharmacon to have minimal targeting of genes in rat cells (see <http://www.dharmacon.com/>).

Antibodies

Rabbit calyntenin-1 and APP C-terminal antibodies were as described previously (15,53). Anti-beta-Actin clone AC-15 and anti-Nicastrin rabbit polyclonal antibodies were from Abcam (Cambridge, MA, USA); anti-ADAM10/Kuz and anti-presenilin-1 N-terminus rabbit polyclonal antibodies were from EMD Chemicals (Gibbstown, NJ, USA); anti-presenilin-1 C-terminus clone PS1-loop, anti-APP mouse clone 22C11 and anti-APP mouse clone 2.F2.19B4 were from Millipore Bioscience Research Reagents (Massachusetts, MA, USA); anti-sAPP α clone 2B3 and anti-sAPP β rabbit polyclonal antibodies were from IBL International (Hamburg, Germany), anti-phospho-threonine-668 APP rabbit polyclonal antibody was from Cell Signaling Technology (Danvers, MA, USA); anti-BACE1 rabbit polyclonal antibody was from Sigma-Aldrich (St Louis, MO, USA); anti-X11 β (Mint 2) and TGN-38 mouse monoclonal antibodies were from BD Biosciences (San Jose, CA, USA); neuron-specific enolase (NSE) antibody was from Dako (Cambridge UK).

Cell culture and transfection

Cortical neurons were obtained from embryonic day 18 rat embryos and cultured in Neurobasal medium containing B27 supplement, 100 IU/ml penicillin, 100 μ g/ml streptomycin and 2 mM L-glutamine (Invitrogen). For siRNA knockdown studies, neurons at DIV4 were treated with 1 μ M siRNAs and analysed at DIV8. For transfection studies, neurons cultured on poly-L-lysine-coated glass cover slips in 12-well plates were transfected using Lipofectamine 2000 (Invitrogen) (1 μ g DNA and 0.5 μ l Lipofectamine 2000) as described by the manufacturer at DIV7 and analysed on DIV8.

A β 1-40 and A β 1-42 ELISAs

Rat/mouse and human A β 1-40 and A β 1-42 ELISAs were obtained from Invitrogen. Samples were processed according to the manufacturer's instructions.

SDS-PAGE and immunoblotting

Cells were harvested for SDS-PAGE by washing with phosphate-buffered saline pre-warmed at 37°C and scraping into SDS-PAGE sample buffer and immediately heating to 100°C. λ -Phosphatase treatment was as described previously (54). Samples were separated on either 8 or 10% (w/v) acrylamide gels or for analyses of APP C-terminal processed fragments on 16% (w/v) acrylamide Tris-tricine gels as described previously (55). Separated proteins were then transferred to Protran nitrocellulose membranes or for Tris-tricine gels, to 0.2 μ m Optitran nitrocellulose membranes (Sigma) using a Transblot system (BioRad) and then processed for immunoblotting. Signals on immunoblots were quantified using ImageJ (National Institutes of Health) after scanning with an Epson Precision V700 Photo scanner essentially as described by us in previous studies (15,56). To ensure the signals obtained were within the linear range, the mean background-corrected optical density (OD) of each signal was interpolated

for an OD calibration curve created using a calibrated OD step tablet (Kodak). Only film exposures that gave OD signals within the linear range of the OD calibration curve were used for the statistical analyses.

Frozen human brain tissue was prepared as a 20% homogenate in ice-cold lysis buffer containing 50 mM Tris-HCl pH 7.4, 0.9% (w/v) NaCl, 0.1% (v/v) Triton X-100, 10 mM sodium fluoride, 1 mM sodium orthovanadate, 2 mM ethylenediamine tetraacetic acid, 1 mM phenylmethylsulfonyl fluoride and complete protease inhibitors (Roche). Following homogenization, samples were centrifuged at 25 000g (av) for 20 min at 4°C and prepared for SDS-PAGE by addition of sample buffer. Signals on immunoblots were simultaneously visualized and the relative amounts quantified using an Odyssey imaging system and associated software (Li-Cor Biosciences, Cambridge, UK). Data were analysed using Student's *t*-test.

Immunofluorescence and time-lapse microscopy

Neurons were prepared for immunostaining by fixing in 4% paraformaldehyde in phosphate-buffered saline (PBS) for 20 min, quenching with 0.05 M NH₄Cl for 15 min, washing in PBS and then permeabilizing with 0.2% Triton X-100 in PBS for 5 min. Following blocking with 5% goat serum (Sigma) in PBS for 30 min, the samples were probed with primary antibodies diluted in blocking solution (PBS containing 5% goat serum). Primary antibodies were then detected using goat anti-mouse and anti-rabbit Igs coupled to Alexa Fluor 488 or 546 (Invitrogen). Confocal images were captured using a Zeiss LSM510Meta confocal microscope equipped with a \times 100 Plan-Apochromat 1.4 N.A. objective (Carl Zeiss Ltd, Welwyn Garden City, UK). Non-confocal images were captured using a Leica DM5000B microscope with \times 63 HCX PL Fluotar phase objective. Co-localization of signals was performed using ImageJ 1.44p with the RG2B co-localization plug-in to determine co-localized pixels. To quantify APP signals in the TGN, TGNs were first marked by immunostaining with TGN-38. APP signals in the delineated areas were then quantified by calculating the average pixel fluorescence intensity using ImageJ. These signals were then normalized by subtracting the background fluorescence signals in each cell which were obtained from measurements in nuclei. Data were analysed using Student's *t*-test.

Live microscopy of APP-EGFP, APP-mCherry and EGFP-calyntenin-1 was performed with an Axiovert S100 microscope (Zeiss) equipped with a Lambda LS Xenon-Arc light source (Sutter Instrument Company, Novato, CA, USA), GFP or GFP-mCherry filter set (Chroma Technology Corp., Rockingham, VT, USA), \times 100 Plan-Apochromat 1.4 N.A. objective (Zeiss), Lambda 10-3 filter wheel (Sutter Instrument Co.) and a Photometrics Cascade-II 512B High Speed EMCCD camera (Photometrics, Tuscon, AZ, USA). Twenty-four hours post-transfection, the cells were transferred to a Ludin imaging chamber (Life Imaging Systems, Basel, Switzerland) mounted on the stage of the microscope. Neurons were maintained in a HEPES-buffered extracellular neuronal solution (composition in mM: NaCl, 140; KCl, 5; NaHCO₃, 5; MgCl₂·6H₂O, 1; CaCl₂, 1.2; Na₂HPO₄, 1.2; Glucose, 10; Hepes, 20; pH 7.4) at 37°C using an objective

Table 1. Case data for human post-mortem brain samples

Stage	Case no.	Sex	Age (years)	Post-mortem delay (h)
Control	A042/01	F	52	44
Control	A278/96	F	77	29
Control	A239/95	F	79	38
Control	A155/95	F	63	34
Control	A094/95	F	80	31
Control	A047/02	F	87	22
Control	A170/00	F	68	9
Control	A322/94	F	62	81
Control	A346/95	M	85	16
Control	A135/95	M	65	24
Control	A134/00	M	86	6
Control	A401/97	M	85	42
Control	A223/96	M	80	11
Control	A133/95	M	85	48
Control	A149/01	M	95	44
Control	A077/00	M	68	53
Control	A066/00	M	61	53
Control	A330/94	M	69	52
Control	A261/94	M	75	85
Braak VI	A157/00	F	75	9
Braak V	A240/06	F	97	12
Braak VI	A010/06	F	67	56
Braak V	A210/05	F	84	18
Braak V–VI	A169/05	F	82	43
Braak V–VI	A168/05	F	84	36
Braak V–VI	A074/05	F	89	29
Braak V	A203/04	F	84	37
Braak V	A221/03	F	81	37
Braak VI	A188/00	F	64	10
Braak VI	A232/00	F	79	8
Braak VI	A074/06	F	69	25
Braak VI	A206/07	M	81	47
Braak V–VI	A186/04	M	71	5
Braak VI	A122/04	M	86	26
Braak VI	A176/01	M	71	41

heater (IntraCell, Royston, UK) and ‘The Box’ Microscope Temperature Control System (Life Imaging Systems). Vesicle movements were recorded for 5 min with a 1 s time-lapse interval using MetaMorph software (Molecular Devices). A movement was defined as a displacement of a particle of at least 2 μm , without reversal of direction of more than 2 μm or pausing for longer than 5 s. For dual imaging of APP-mCherry and EGFP-calsyntenin-1 transport, co-transfected neurons were imaged using a two-channel simultaneous imaging system (Dual View, Photometrics) and vesicle movements were recorded for 2 min with a 2 s time-lapse interval using MetaMorph software (Molecular Devices). Image analysis was performed with ImageJ. The flux rates for APP-EGFP transport in both anterograde and retrograde directions were determined essentially as described (8,27) by counting the numbers of APP-EGFP particles that crossed a defined line in mid axons per minute. Statistical significance was determined using Student’s *t*-test. Images presented were constructed using Adobe CS5 and have been smoothed in ImageJ.

Human tissues

The post-mortem human frontal cortex was obtained from control and pathologically confirmed cases of Alzheimer’s

disease (Braak Stage V–VI) (Table 1). All tissue collection and processing were carried out under the regulations and licensing of the Human Tissue Authority and in accordance with the Human Tissue Act, 2004. Calsyntenin-1 signals were normalized against NSE, as neuronal loss or astrogliosis may bias the data toward downregulation of neuronal proteins.

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