

The novel cargo Alcadein induces vesicle association of kinesin-1 motor components and activates axonal transport

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Alcadein α (Alc α) is an evolutionarily conserved type I membrane protein expressed in neurons. We show here that Alc α strongly associates with kinesin light chain ($K_D \approx 4\text{--}8 \times 10^{-9}$ M) through a novel tryptophan- and aspartic acid-containing sequence. Alc α can induce kinesin-1 association with vesicles and functions as a novel cargo in axonal anterograde transport. JNK-interacting protein 1 (JIP1), an adaptor protein for kinesin-1, perturbs the transport of Alc α , and the kinesin-1 motor complex dissociates from Alc α -containing vesicles in a JIP1 concentration-dependent manner. Alc α -containing vesicles were transported with a velocity different from that of amyloid β -protein precursor (APP)-containing vesicles, which are transported by the same kinesin-1 motor. Alc α - and APP-containing vesicles comprised mostly separate populations in axons *in vivo*. Interactions of Alc α with kinesin-1 blocked transport of APP-containing vesicles and increased β -amyloid generation. Inappropriate interactions of Alc- and APP-containing vesicles with kinesin-1 may promote aberrant APP metabolism in Alzheimer's disease.

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Introduction

Conventional kinesin (kinesin-1) is composed of two heavy (KHC) and two light (KLC) chains (Hirokawa, 1998). The N-terminal half of KLC interacts with KHC, and the

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C-terminus of KLC links to membrane cargo either directly or via cytosolic adaptor proteins such as the JNK-interacting protein 1 (JIP1) (Gauger and Goldstein, 1993; Gindhart and Goldstein, 1996; Verhey *et al.*, 2001). The combinations of cargoes, adaptor proteins, and motor molecules contribute to the diversity of vesicular transport systems (Hirokawa, 1998), the dysfunction of which may underlie some neurodegenerative disorders (Guzik and Goldstein, 2004) such as Alzheimer's disease (AD) (Stokin *et al.*, 2005).

Amyloid β -protein precursor (APP) has been implicated in the development and progression of AD (Gandy, 2005) and is proposed to be a kinesin-1 cargo. APP can bind to kinesin-1 directly (Kamal *et al.*, 2000; Gunawardena and Goldstein, 2001) or through JIP1 (Inomata *et al.*, 2003; Matsuda *et al.*, 2003). Although the cargo hypothesis has been challenged (Lazarov *et al.*, 2005) and it is unclear how APP functions mechanistically in vesicle transport, disruption of APP-containing vesicle transport may alter APP metabolism, thereby increasing levels of neurotoxic β -amyloid ($A\beta$) (Gunawardena and Goldstein, 2001; Stokin *et al.*, 2005).

Alcadein (Alc) is an evolutionarily conserved type I membrane protein, which has four isoforms, Alc α 1, Alc α 2, Alc β , and Alc γ , in mammals (Araki *et al.*, 2003). These proteins were isolated independently as postsynaptic Ca^{2+} -binding proteins called the calsynenins (Vogt *et al.*, 2001). Alc associates with APP in neurons through cytoplasmic interactions of both proteins with X11-like (X11L) (Araki *et al.*, 2003), a neuron-specific adaptor protein (Tomita *et al.*, 1999). Formation of a tripartite APP/X11L/Alc complex stabilizes both APP and Alc proteins metabolically. Dissociation of X11L from the complex induces coordinated cleavages of APP and Alc, generating the $A\beta$ fragment from APP and the β -Alc peptide from Alc, in addition to the release of their respective cytoplasmic-domain fragments, AICD and AlcICD (Araki *et al.*, 2004). Thus, Alc may be similar to APP in function as well as metabolism.

Here, we present evidence that Alc is a kinesin-1 cargo and exhibits novel functions. The transport of Alc-containing vesicles competes with that of APP-containing vesicles for kinesin-1, and disruption of transport of APP-containing vesicles increases $A\beta$ generation. This analysis contributes to our understanding of both the pathogenesis of AD and the physiological function of Alc.

Results

Direct interaction of Alc with KLC

A yeast two-hybrid screen using the cytoplasmic domain of Alc α (amino acids 872–969 of mouse Alc α 1) as bait isolated 11 full-length and partial cDNA clones encoding KLC1 (see Figure 1A and Supplementary Figure S1 for the schematic structures of KLC1 encoded by the isolated cDNA clones; Supplementary data is shown in Sup_1.pdf.) and one partial

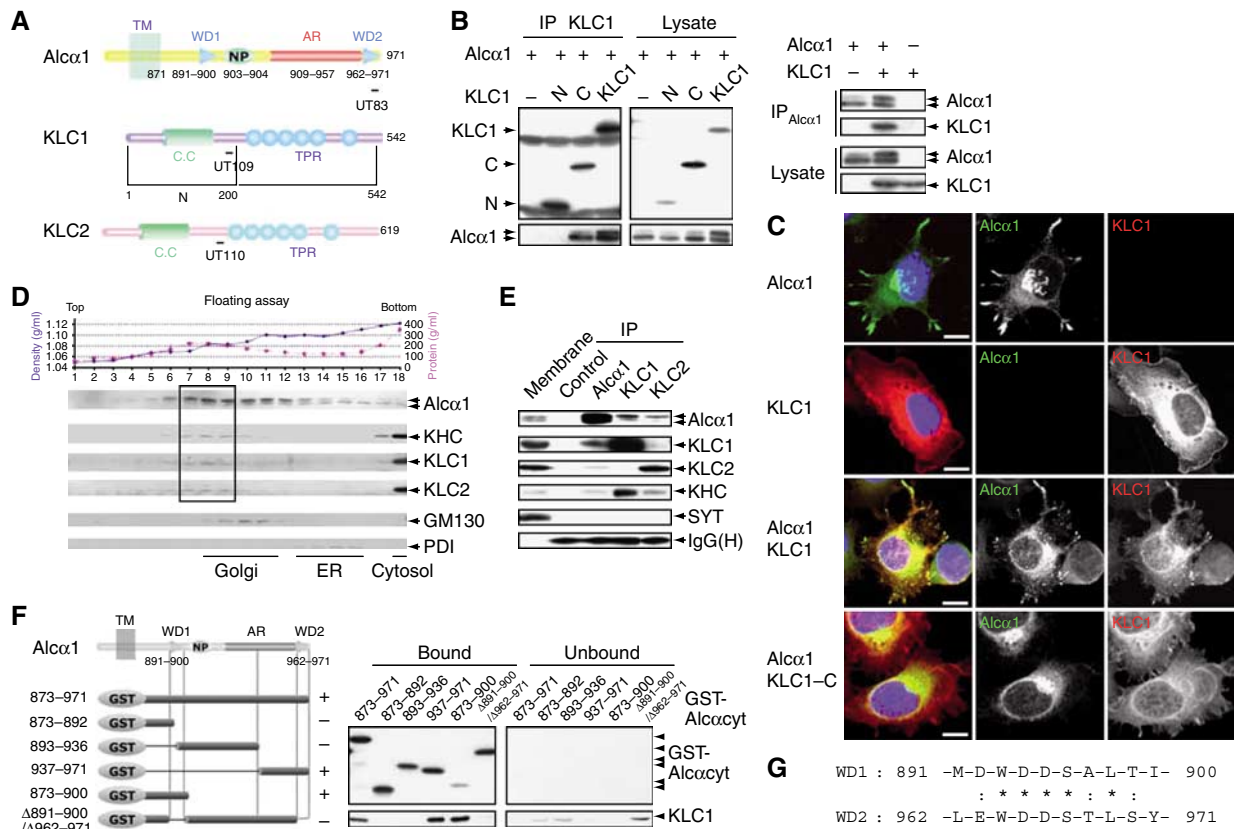


Figure 1 Interaction of Alc α with KLC and determination of the KLC-binding site on Alc α . (A) Structure of the cytoplasmic domains of Alc α , KLC1, KLC1-N (N), KLC1-C (C), and KLC2. Numbers represent amino-acid sequences. TM, transmembrane domain; WD, tryptophan- and aspartic acid-containing sequence (see 'G'); NP, Asn-Pro motif; AR, acidic region; CC, coiled-coil domain; TPR, tetratricopeptide repeat. Epitopes for the UT83, UT109, and UT110 antibodies are indicated with bars. (B) (Left panel) Co-immunoprecipitation of Alc α 1 with amino-terminal FLAG-tagged KLC1 and deletion constructs. HEK293 cells were transiently cotransfected with pcDNA3-hAlc α 1 and pcDNA3.1-FLAG-mKLC1 (KLC1), pcDNA3.1-FLAG-mKLC1-N (N), pcDNA3.1-FLAG-mKLC1-C (C), or pcDNA3.1-FLAG (-). Cell lysates were immunoprecipitated with an anti-FLAG (M2) antibody. The immunoprecipitates (IP) and lysate were analyzed by Western blotting with M2 and UT83. (Right panel) Co-immunoprecipitation of KLC1 with Alc α 1. HEK293 cells were transiently transfected with pcDNA3-hAlc α 1 and pcDNA3.1-mKLC1. Transfection with plasmid (+) or vector alone (-) is indicated. Cell lysates were immunoprecipitated with an anti-Alc α (UT83) antibody. The immunoprecipitates (IP) and lysate were analyzed by Western blotting with UT83 and UT109. (C) Localization of Alc α 1 and KLC1. CAD cells were transiently cotransfected with the indicated combinations of pcDNA3-hAlc α 1, pcDNA3.1-FLAG-mKLC1, and pcDNA3.1-FLAG-mKLC1-C for 48 h and differentiated for 24 h by depleting serum. Alc α 1 and FLAG-tagged KLC1 were immunostained with UT83 (green) and M2 (red) antibodies, respectively. Nuclei were stained using 4',6'-diamidino-2-phenylindole dihydrochloride (DAPI, blue). Merged signals are shown at the left. Scale bar, 5 μ m. (D) Membrane fractionation of brain tissues. The post-nuclear supernatant of adult mouse brains was fractionated by 0–28% iodixanol density gradient centrifugation (Araki *et al*, 2003). (Upper panel) The density (blue circles) and protein concentration (pink squares) are indicated. Fraction numbers are indicated along the abscissa. (Lower panel) The fractions were analyzed by Western blotting with antibodies to Alc α (UT83), KHC (H2), KLC1 (UT109), KLC2 (UT110), and the Golgi-resident GM130 (clone no. 35) and the ER-resident PDI (1D3). The dotted square indicates the fractions containing the highest levels of Alc α , KHC, KLC1, and KLC2. (E) Co-immunoprecipitation of Alc α with KLC. The mouse brain membrane fraction (500 μ g protein) was solubilized, and Alc α 1, KLC1, and KLC2 were immunoprecipitated with the UT83, UT109, and UT110 antibodies, respectively, or a control non-immune antibody. The membrane fraction (Membrane, 10 μ g protein) and immunoprecipitates (IP) were analyzed by Western blotting with antibodies specific for Alc α (UT83), KLC1 (UT109), KLC2 (UT110), KHC (H2), and SYT (clone no. 41). IgG(H) indicates the IgG heavy chain (rabbit). (F) Determination of the KLC-binding site on Alc α . (Left) Various GST-fusion proteins of the Alc α cytoplasmic domain (shown schematically, numbers are amino-acid residues) were incubated with HEK293 cell lysates expressing FLAG-KLC1. FLAG-KLC1 bound to GST-Alc α cyt was recovered using glutathione beads. (Right) The bound and unbound FLAG-KLC1 were analyzed by Western blotting with the M2 antibody. GST-Alc α 1 protein constructs were detected by Western blotting with an anti-GST antibody. In the left panel, '+' indicates positive binding and '-' indicates negative binding. (G) Amino-acid sequences of WD1 and WD2 of Alc α 1. Numbers indicate amino-acid residues. Identical (*) and similar (:) amino-acid residues are indicated.

cDNA clone encoding KLC2. The interaction between Alc α 1 and KLC1 was confirmed by co-immunoprecipitation in HEK293 cells overexpressing Alc α 1 and KLC1 (Figure 1A and B, right). Immunoprecipitates obtained with an anti-Alc α antibody contained KLC1, indicating that Alc α interacts with KLC1. To identify the Alc α -binding domain, FLAG-tagged amino- (KLC1-N or N) and carboxy- (KLC1-C or C) terminal halves of KLC1 (Figure 1A) were overexpressed. An anti-FLAG antibody co-immunoprecipitated Alc α 1 with KLC1

and with KLC1-C, but not with KLC1-N (Figure 1B, left). These data indicate that Alc α interacts with the KLC1 carboxy-terminus, which includes the tetratricopeptide repeat (TPR; see Figure 1A), with which other KLC-binding proteins interact (Gindhart and Goldstein, 1996). Alc α 1 was also co-immunoprecipitated with FLAG-KLC2 (data not shown).

Alc α 1 appears as two bands on SDS-PAGE gels. The more slowly migrating band increased in intensity in the presence of KLC1 and had a mobility similar to that of brain Alc α

(Figure 1B and Supplementary Figure S2). HEK293 cells express substantial amounts of endogenous KHC and KLC2 but lower levels of endogenous KLC1 (Supplementary Figure S2A), as do undifferentiated CAD cells (data not shown), suggesting that the expression of exogenous KLC1 may increase the formation of functional kinesin-1 in both cell types because increased number of transporting Alcx vesicles were observed (data not shown for HEK293 cells and Figure 1C for CAD cells). We determined that Alcx1 was post-translationally modified with both high-mannose (arrow 2) and complex (arrow 1) N-glycans (Supplementary Figure S2B). The predominance of the slow-migrating Alcx1 increased in the presence of KLC1 but not KLC1-C (Figure 1B, left), which does not bind to KHC (Gauger and Goldstein, 1993), suggesting that trafficking of Alcx-containing vesicles by kinesin-1 facilitates complex N-glycosylation.

Alcx colocalized with KLC1 in CAD cells (Figure 1C). CAD cells overexpressing Alcx1 in the presence (Figure 1C, third rows) or absence (first rows) of KLC overexpression began to extend processes that were not observed in cells not expressing Alcx1 (second rows) after serum depletion. Alcx1 protein localized to neurite tips, as well as the ER and Golgi (third rows), whereas FLAG-KLC1 was located in the cytoplasm (second rows). A few small vesicles were observed in cells expressing Alcx1 alone (first rows), but a large number of small vesicles appeared when Alcx1 was coexpressed with FLAG-KLC1, and the proteins were colocalized (third rows), suggesting that Alcx binds the active kinesin-1 motor complex and uses it to transport vesicles to neurite tips. Such vesicles were not observed in cells coexpressing KLC1-C with Alcx1 (fourth rows), as KLC1-C is not able to bind KHC.

Alcx was included in a protein complex containing KLC and KHC *in vivo*. Mouse brain membranes were fractionated by iodixanol density gradient centrifugation, and Alcx, KHC, KLC1, KLC2, the Golgi-resident protein GM130, and the ER-resident protein disulfide isomerase (PDI) were detected in these membranes (Figure 1D). The complex N-glycosylated Alcx was recovered with kinesin-1 subunits from vesicle fractions that were less dense than the Golgi membrane fractions. The faster-migrating high-mannose N-glycosylated Alcx was largely recovered in the ER. The membranes resulted by 100 000 g centrifugation were solubilized, immunoprecipitated with anti-Alcx, anti-KLC1, anti-KLC2, or non-immune control antibodies, and subjected to Western blot analysis (Figure 1E). Alcx, KLC1, KLC2, and KHC were immunoprecipitated by anti-Alcx, anti-KLC1, as well as the anti-KLC2 antibody. As a control, synaptotagmin (SYT) was not co-immunoprecipitated by any of the antibodies. These results suggest that Alcx associates with kinesin-1 *in vivo*.

A pull-down assay with recombinant KLC1 and a GST-Alcx cytoplasmic domain fusion protein (GST-Alcx_{cyt}) demonstrated a direct interaction between KLC1 and Alcx (Supplementary Figure S3A). KLC1 bound to GST-Alcx_{cyt} with a dissociation constant (K_D) of $\approx 7.9 \times 10^{-9}$ M. Surface plasmon resonance (SPR) analysis also demonstrated a strong affinity ($K_D \approx 4.3 \times 10^{-9}$ M) between GST-Alcx_{cyt} and KLC1: the interaction had an association rate (K_{ass}) of approximately 1.3×10^3 M/s and a dissociation rate (K_{diss}) of approximately 5.5×10^{-6} M/s (Supplementary Figure S3B). This low dissociation rate indicates a very strong binding between Alcx and KLC1.

The region of Alcx that binds KLC1 was determined by a pull-down assay with glutathione beads bearing GST fusion

proteins (Figure 1F). KLC bound to the Alcx fusion protein containing amino acids 873–971, but did not bind to fusion proteins lacking both WD motifs, WD1 = amino acids 891–900 and WD2 = amino acids 962–971. Each WD motif contains the consensus, D/E-W-D-D-S-A/T-L-T/S (Figure 1G). Deletion of either WD1 or WD2 preserved KLC1 binding, indicating that one WD motif is sufficient for KLC binding. KLC binding to the WD motifs was completely abolished when Ala was substituted for the conserved amino-acid residues in the WD motifs (unpublished observations and Figure 6A), demonstrating a highly specific interaction between Alcx and KLC. These findings *in vitro* strongly suggest that Alcx1 can function as a kinesin-1 cargo.

Alcx is a novel cargo for kinesin-1 and can mediate the vesicle association of kinesin-1 motor components

To investigate the characteristics of Alcx1 as a cargo, an Alcx1-GFP fusion protein was expressed in differentiating CAD cells and assayed for vesicular transport. Fast anterograde transport of Alcx1 cargo was observed, together with a minor amount of retrograde movement (Figure 2A, panel 1, and Supplementary Movie 1, part 1 in Sup_2.mov). Approximately 80% of the total Alcx1 cargo vesicles (375 vesicles in 25 cells) showed anterograde transport and 10–15% showed retrograde movement (Figure 2A, panel 2). The anterograde vesicles were transported at a velocity of 1.48 ± 0.45 $\mu\text{m/s}$ (mean \pm s.d.), which was calculated from the movement of 4125 vesicles (Figure 2A, panel 3) and was consistent with the velocity of 1.32 ± 0.58 $\mu\text{m/s}$ observed in primary cultured mouse cortical neurons expressing Alcx1-Venus (Supplementary Figure S4A and Movie 2, part 1 in Sup_3.mov) and with the velocity of 1.47 ± 0.76 $\mu\text{m/s}$ observed in non-neuronal HEK293 cells expressing Alcx1-Venus and KLC1 (Supplementary Movie 3, part 1 in Sup_4.mov). Alcx1-Venus transport was suppressed by treatment with nocodazole and Alcx1-mRFP-containing vesicles moved on microtubule tracks composed of YFP-tubulin in HEK293 cells (Supplementary Movie 3, parts 2 and 3 in Sup_4.mov). These observations indicate that Alcx is transported on microtubule tracks.

To determine whether transport required kinesin-1, endogenous KLC1 and KLC2 expression was knocked down. Differentiating CAD cells expressing siRNA directed against KLC1 or KLC2 showed almost no expression of the respective KLC (data not shown). When siRNA treatment against both KLC1 and KLC2 was performed in differentiating CAD cells, fast anterograde transport of vesicles containing Alcx1-GFP was dramatically inhibited, whereas retrograde transport was largely unaffected (Figure 2B, panel 1, and Supplementary Movie 1, part 2 in Sup_2.mov). Approximately 60% of vesicles (375 vesicles in 25 cells) were stationary (Figure 2B, panels 2 and 3), which is likely to result from interruption of anterograde transport owing to lack of KLC. Identical results were obtained in primary cultured mouse cortical neurons (Supplementary Figure S4B and Movie 2, part 2 in Sup_3.mov).

Interestingly, Alcx showed a striking effect on the vesicle association of kinesin-1 motor components. GFP-KLC1 was expressed in differentiating CAD cells in the presence or absence of Alcx1 expression. GFP-KLC alone showed a diffuse distribution, as described previously (Verhey *et al*, 1998), which also agreed with KLC1 immunostaining

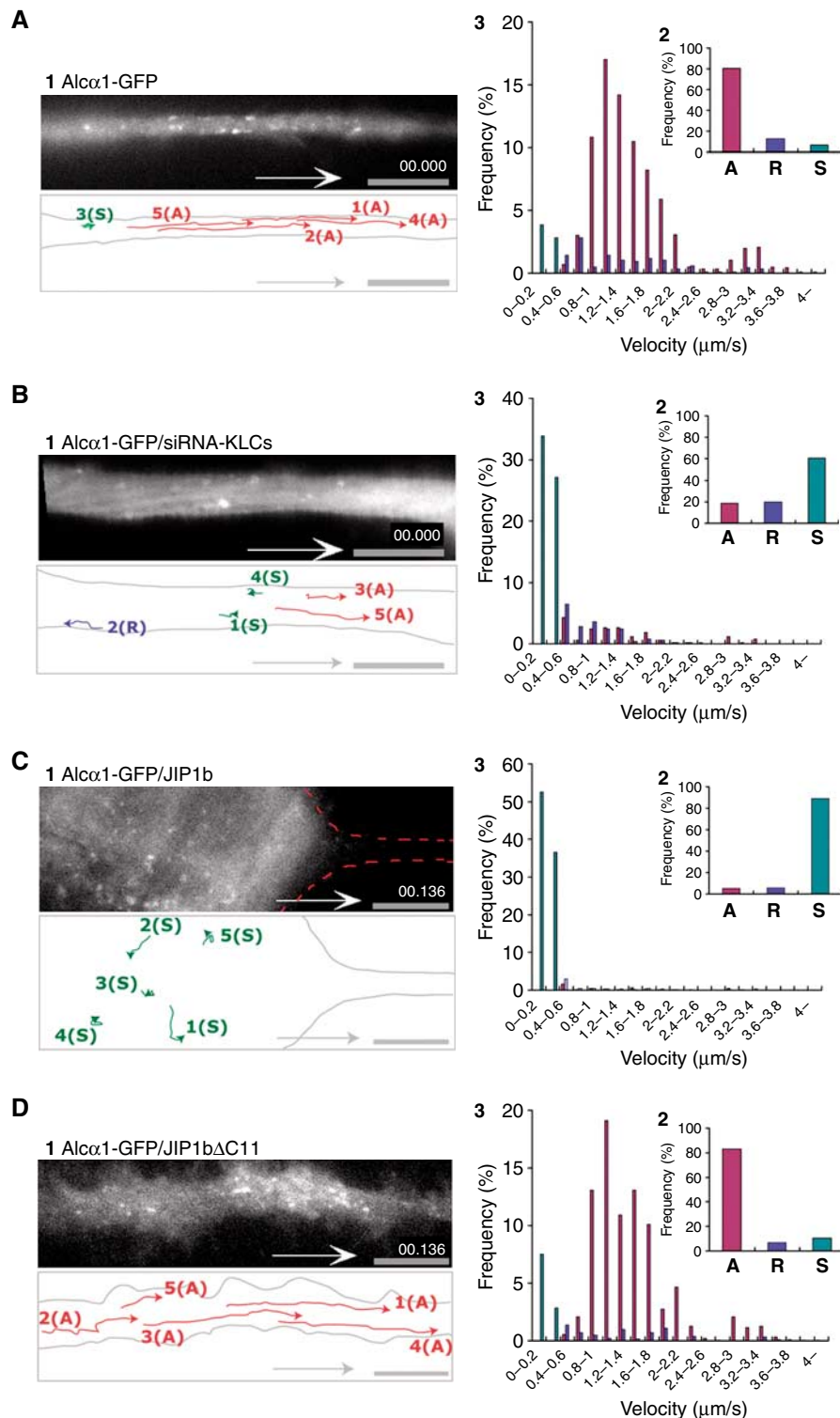


Figure 2 Kinesin-1-dependent anterograde transport of Alc α cargo in living neuronal cells. (A, B) Anterograde movement of Alc α cargo vesicles in an axon. (A) Differentiating CAD cells expressing Alc α 1-GFP were observed using TIRF microscopy (panel 1). (B) Kinesin-1-dependent transport of Alc α 1 cargo in neuronal cells. Differentiating CAD cells expressing Alc α 1-GFP in which KLC1 and KLC2 expression has been knocked down using siRNA were observed using TIRF microscopy (panel 1). Vesicle movements in the dotted square were tracked with time-lapse imaging and are indicated with colored lines and numbers (see Supplementary Movie 1, parts 1 and 2 in Sup_2.mov). Scale bar, 5 μm . (C, D) Inhibition of anterograde transport of Alc α 1 cargo by expression of JIP1b. Differentiating CAD cells expressing Alc α 1-GFP in the presence of JIP1b (C) and JIP1b Δ C11 (D) were observed using TIRF microscopy (panel 1). Vesicle movements in the dotted square were tracked with time-lapse imaging and are indicated with colored lines and numbers (see Supplementary Movie 1, parts 3 and 4 in Sup_2.mov). Scale bar, 5 μm . (A–D) Red lines indicate tracks of anterograde transport, blue lines indicate tracks of retrograde transport, and green spots indicate stationary vesicles moving at less than 0.4 $\mu\text{m/s}$ (panel 1). Alc α 1 and APP cargo vesicles transported anterogradely ('A') and retrogradely ('R'), and stationary vesicles ('S') in 25 cells were counted with Metamorph software and the fraction of the total number of vesicles (%) is indicated (panel 2). Distribution (%) of anterograde (red) and retrograde (blue) transport velocity of Alc α 1 cargo and of stationary vesicles (green) is shown (panel 3).

(Figure 1C). In contrast, in the presence of Alcα1-, KLC1-containing vesicles were visible and underwent anterograde transport (Figure 3A and B, panels 1 and 2, and Supplementary Movie 4, parts 1 and 2 in Sup_5.mov). The velocity of GFP-KLC1 in anterograde transport was distributed among two populations of vesicles moving at 1.2–1.4 μm/s or ~3.0 μm/s (Figure 3B, panel 3). These values are consistent with the velocities of Alcα1- and APP cargo-containing vesicles, respectively. Faster moving vesicles at ~3.0 μm/s may be transported with APP/JIP1b or another cargo following the vesicle association of kinesin-1 component by Alcα1. When GFP-KHC was expressed, vesicles containing GFP-KHC were detected and underwent anterograde transport in the absence of exogenously expressed Alcα1 (Figure 3C and Supplementary Movie 4, part 3 in Sup_5.mov). The velocity of GFP-KHC agreed with that of Alcα1 (~1.5 μm/s) and APP (~3 μm/s), indicating that kinesin-1 composed of GFP-KHC and endogenous KLC was associated to membrane via endogenous APP/JIP1b and Alcα1 (Figure 3C, panel 3). The vesicle association of GFP-KHC was suppressed by coexpression of unlabeled KLC (Figure 3D and Supplementary Movie 4, part 4 in Sup_5.mov), consistent with the previous observation that overexpression of KLC inhibits binding of KHC to microtubules (Verhey *et al*, 1998). However, GFP-KHC-containing vesicles were again visible and underwent anterograde transport when Alcα1 was coexpressed in the KLC-overexpressing cells (Figure 3E and Supplementary Movie 4, part 5 in Sup_5.mov). This result cannot be simply due to the quenching and/or masking of overexpressed KLC1 by Alcα1, as overexpression of APP or JIP1b does not cause GFP-KHC to reassociate with vesicles under the same conditions (Figure 3F and Supplementary Movie 4, part 6 in Sup_5.mov). JIP1b and APP did not show any ability to induce vesicle association of GFP-KLC (Figure 3G and Supplementary Movie 4, part 7 in Sup_5.mov). These findings indicate that Alcα1 modulates the vesicle association of kinesin-1 motor components in axonal cargo transport.

Alc blocks transport of APP-containing vesicles and JIP1b blocks transport of Alc cargo vesicles

APP is transported by the kinesin-1 motor system (Kaether *et al*, 2000; Kamal *et al*, 2000; Gunawardena and Goldstein, 2001). Transport of APP-containing vesicles by kinesin-1 was observed by movement of APP-GFP in differentiating CAD cells (Figure 4A, panel 1, and Supplementary Movie 5, part 1 in Sup_6.mov). The anterograde transport velocity of APP-containing vesicles was extremely fast; most vesicles moved at a speed of 3.0–3.6 μm/s with an average of 3.03 ± 1.07 μm/s (Figure 4A, panel 3). Transport of APP-containing vesicles was suppressed when endogenous KLC1 and KLC2 expression

was knocked down (Figure 4B and Supplementary Movie 5, part 2 in Sup_6.mov), suggesting that kinesin-1 transports APP cargoes at a velocity distinct from that of Alcα1 cargoes.

Anterograde transport of APP-GFP-containing vesicles was suppressed in the presence of Alcα1 or its cytoplasmic domain fragment Alcα1CD, a physiologically produced peptide generated by Alcα1 processing (Araki *et al*, 2004). However, the mutant Alcα1CD (AWAA), in which the KLC-interacting sequence D/E-W-D-D-S within the WD1 and WD2 motifs is altered to A-W-A-A-S making the mutant unable to bind KLC, failed to inhibit the transport of APP-containing vesicles (Figure 4C–E and Supplementary Movie 5, parts 3–5). These observations suggest that APP-containing vesicles are transported in a kinesin-1-dependent manner and that Alcα1 cargo vesicles compete with APP-containing vesicles for kinesin-1.

Conversely, movement of Alcα1 cargo vesicles was suppressed in differentiating CAD cells following expression of JIP1b. Under these conditions, axonal transport of Alcα1 cargo was not observed and vesicles remained in the cell body (Figure 2C and Supplementary Movie 1, part 3 in Sup_2.mov). Expression of JIP1bΔC11, which lacks the KLC-binding site, did not inhibit axonal transport of Alcα1 cargo (Figure 2D and Supplementary Movie 1, part 4 in Sup_2.mov), indicating that the interaction of JIP1b with KLC1 may compete with KLC1 for Alcα1.

To analyze the role of JIP1b in cargo selection, we first asked whether JIP1b interacts with Alcα1. Alcα1 was coexpressed in HEK293 cells with known FLAG-tagged Alcα-binding proteins (X11L and FE65) (Araki *et al*, 2003, 2004) or with FLAG-JIP1b and FLAG-KLC1, and the lysates were immunoprecipitated using an anti-FLAG antibody. Alcα1 coimmunoprecipitated with X11L, FE65, and KLC1, but not JIP1b (Figure 5A). This result indicates that Alcα1 does not associate with KLC via a JIP1b bridge. Lysates of HEK293 cells overexpressing KLC1, Alcα1, and JIP1b were immunoprecipitated with an anti-Alcα antibody. The interaction between Alcα1 and KLC1 was inhibited by JIP1b in a dose-dependent manner (Figure 5B, left), but not by JIP1bΔC11 (Figure 5B, right). These results indicate that JIP1b binds to KLC1 but not Alcα and suggest that both Alcα and JIP1b share the kinesin-1 motor and that the KLC of kinesin-1 selects an Alcα cargo in the absence of JIP1b.

APP-containing vesicles are transported with a velocity different from that of Alcα cargo vesicles (Figures 2 and 4). This is not a result derived from transport inhibition that the transport of one class of vesicle and/or organelle leads to a traffic jam that blocks the movement of other vesicles and/or organelles, because APP-GFP and Alcα1-mRFP are transported in the same cell at different velocities without a trafficking jam in our experimental condition and the transport of both proteins was suppressed in the same cell when

Figure 4 Transport of APP-containing vesicles and suppression of transport by expression of Alcα1 and Alcα1CD in living neuronal cells. (A) Anterograde transport of APP-containing vesicles in an axon. Differentiating CAD cells expressing APP-GFP were observed using TIRF microscopy (panel 1). (B) Kinesin-1-dependent transport of APP-containing vesicles in neuronal cells. Differentiating CAD cells expressing APP-GFP and in which KLC1 and KLC2 expression has been knocked down using siRNA were observed using TIRF microscopy (panel 1). (C–E) Anterograde transport of APP-containing vesicles in differentiating CAD cells expressing APP-GFP in the presence or absence of Alcα1 or Alcα1CD. Axonal transport of APP-GFP in the presence of Alcα1 (C), Alcα1CD (D), or Alcα1CD(AWAA) (E) was observed with TIRF microscopy. Vesicle movements in the dotted square were tracked with time-lapse imaging and are indicated with colored lines and numbers (see Supplementary Movie 5, in Sup_6.mov). Red lines indicate vesicles transported anterogradely ('A'), blue lines indicate vesicles transported retrogradely ('R'), and green spots indicate stationary vesicles moving at less than 0.5 μm/s ('S'). Scale bar, 5 μm. The vesicles in 25 cells were counted with Metamorph software and the fraction of the total number of vesicles (%) is indicated (panel 2). Distribution (%) of anterograde (red) and retrograde (blue) transport velocity of APP-containing vesicles and of stationary vesicles (green) is indicated (panel 3).

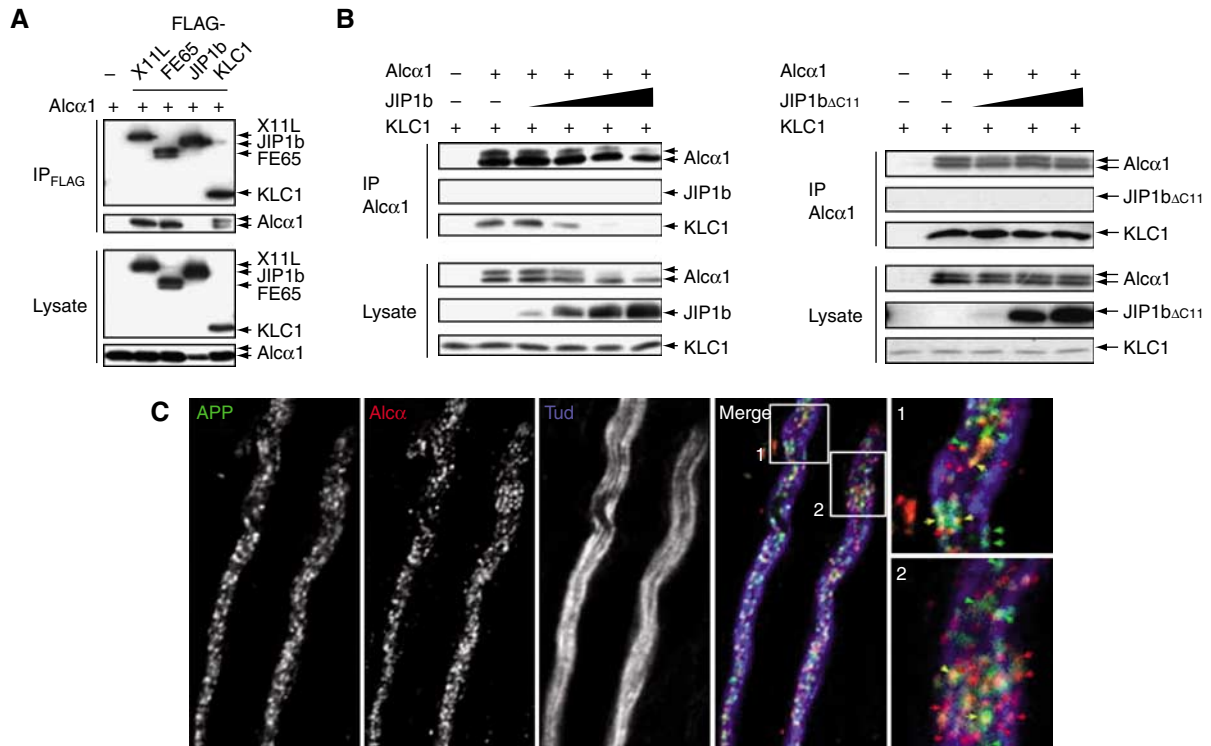


Figure 5 Separate transport of APP- and Alcα-containing vesicles by kinesin-1. (A) Interaction of Alcα1 with adaptor proteins. HEK293 cells were transiently transfected with pcDNA3-hAlcα1 (+) in the presence of pcDNA3-FLAG-hX11L, pcDNA3-N-FLAG-FE65, pcDNA3-FLAG-JIP1b, pcDNA3.1-FLAG-mKLC1, or vector alone (–). Cells were lysed and immunoprecipitated with the anti-FLAG M2 antibody. Immunoprecipitates (IP) and cell lysates (Lysate) were analyzed by Western blotting with the M2 and anti-Alcα UT83 antibodies. (B) Effect of wild-type and mutant JIP1b on the Alcα1-KLC1 interaction. (Left) HEK293 cells were transiently cotransfected with pcDNA3.1-mKLC1 (0.5 μg), pcDNA3-hAlcα1 (0.5 μg), and 0, 0.05, 0.2, 0.7, or 2.2 μg pcDNA3-JIP1b. Cell lysates were immunoprecipitated with an anti-Alcα (UT83) antibody and immunoprecipitates (IP) and cell lysates (Lysate, 10 μg protein) were analyzed by Western blotting with the UT83, UT72, and UT109 antibodies. (Right) Cells were similarly cotransfected using 0, 0.2, 0.7, or 2.2 μg pcDNA3.1-JIP1bΔC11. The UT83 immunoprecipitate (IP) and cell lysate (Lysate, 10 μg protein) were analyzed as described above. Transfection with 0.5 μg plasmid (+) or vector alone (–) is indicated. (C) APP- and Alcα-containing vesicles in adult mouse sciatic nerve. Isolated axons of sciatic nerves were immunostained with anti-APP (αAPP/C, green), anti-Alcα (co190, red), and anti-neuronal Class III β-tubulin (TUJ1, blue; showing axons) antibodies. Signals are merged and magnified in the lower panels. Yellow arrows indicate vesicles containing both APP and Alcα. Scale bar, 5 μm.

the expression of KLCs was knocked down (Supplementary Figure S5 and Movie 6 in Sup_7.mov). However, it needs to be confirmed whether APP-containing vesicles and Alcα cargos are independently transported *in vivo*. To examine this, we localized endogenous APP- and Alcα-containing vesicles in the sciatic nerve of adult mice (Figure 5C). Most APP (green) and Alcα (red) axonal vesicles were separate, but a few stained for both proteins. We confirmed this in a separate study by staining with the independent synaptic vesicle marker, synaptophysin (Supplementary Figure S6). APP and Alcα were colocalized on almost 30% of the vesicles, whereas both Alcα and APP localized independently of synaptophysin. These results indicate that APP and Alcα are largely transported separately, although a minor population of vesicles may contain both proteins. Alternatively, kinesin-1 may exchange APP-containing vesicles for Alcα cargo vesicles, or vice versa, at the points of colocalization.

Suppressed transport of APP-containing vesicles by Alcα facilitates Aβ generation

A previous report indicated that impaired axonal transport resulting from a reduction in the levels of kinesin-1 increased Aβ levels (Stokin *et al*, 2005). Therefore, we hypothesized that the relative amounts of JIP1b and Alcα, which facilitate

or retard APP transport, respectively, may be important for the regulation of Aβ generation. When Alcα cargoes predominate, the movement of APP-containing vesicles is severely impaired (Figure 4C and Supplementary Movie 5, part 3 in Sup_6.mov). We examined Aβ generation in cells in which APP-containing vesicles are being actively transported by association with kinesin-1 in the presence of JIP1b, or in which transportation of those vesicles is retarded due to overexpression of AlcαICD. AlcαICD was used in the following study instead of Alcα1, as the overexpression of Alcα1 competitively inhibits the cleavage of APP, resulting in an apparent reduction of Aβ generation. AlcαICD directly associates with KLC1 and suppresses the transport of APP as well as Alcα1 (Figure 4D and Supplementary Movie 5, part 4 in Sup_6.mov), but does not compete with APP for cleavage (Supplementary Figure S7).

When KLC1, APP, and JIP1b were overexpressed in HEK293 cells and cell lysates were immunoprecipitated with anti-KLC1 antibody, JIP1b and APP co-immunoprecipitated with KLC1, but their recovery was decreased by co-expression of AlcαICD(WT) (Figure 6A). This effect was also observed when the Asn-Pro motif of AlcαICD was altered to Ala-Ala (NPAA), which blocks the ability of Alcα to bind to X11L and FE65 (Araki *et al*, 2004), but not to KLC (Figure 1F).

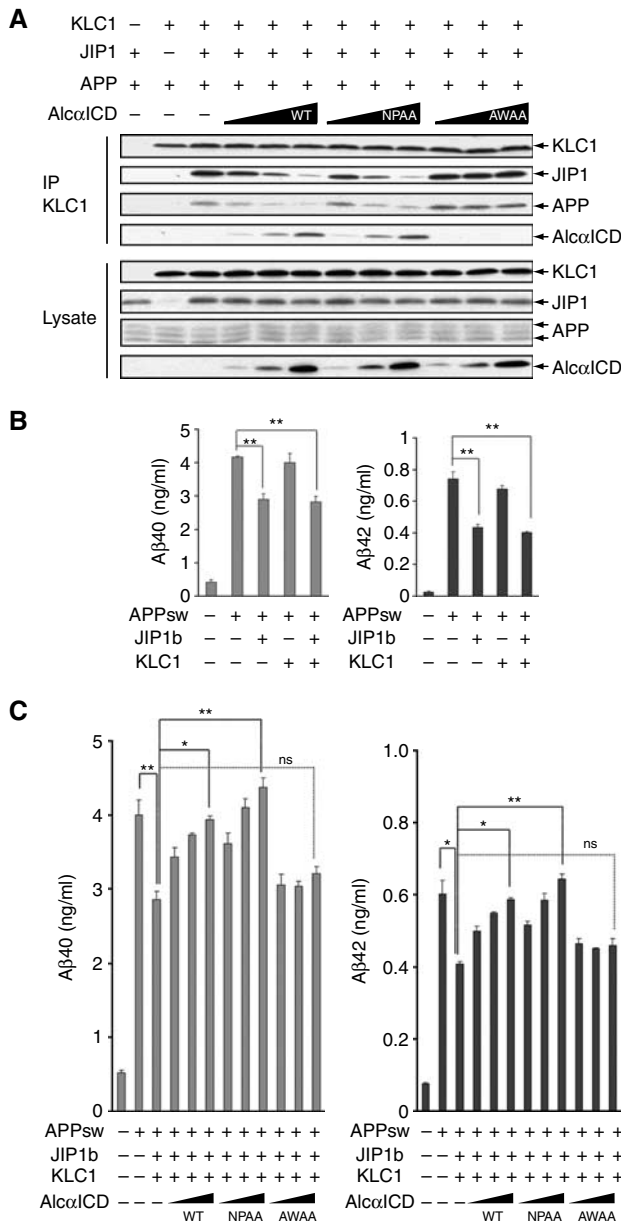


Figure 6 Increased A β production by suppressed transport of APP-containing vesicles. (A) Alc α ICD expression interferes with JIP1b-mediated association of APP and KLC1. HEK293 cells were transiently cotransfected with pcDNA3.1-mKLC1, pcDNA3-JIP1b, pcDNA3-hAPP695, and 0, 0.25, 0.5, or 1.0 μ g pcDNA3-hAlc α ICD (WT), pcDNA3-hAlc α ICD/NPAA (NPAA), or pcDNA3-hAlc α ICD/AWAA (AWAA). Transfection with 0.5 μ g plasmid (+) and vector alone (-) is indicated. Cell lysates were subjected to immunoprecipitation with an anti-KLC1 antibody (UT109), and the immunoprecipitates (IP) and cell lysates (Lysate, 10 μ g protein) were analyzed by Western blotting with UT109, anti-JIP1b (UT72), anti-APP (α APP/c; Sigma), and anti-Alc α (UT83) antibodies. (B) A β secretion from APPsw mutant HEK293 cells expressing JIP1b or KLC1 alone. A β secretion from HEK293 cells used in (A) in which transfection of pcDNA3-hAPP695sw was substituted for pcDNA3-hAPP695. At 48 h after transfection, the culture medium was collected and analyzed for A β 40 (left) and A β 42 (right) levels using a sandwich ELISA (Araki *et al*, 2003). Concentrations of A β 40 and A β 42 are presented as means \pm s.e. ($n = 3$). (C) HEK293 cells transfected with pcDNA3-hAPP695sw and the indicated plasmids, as described in (A), were used for A β assay. A β 40 (left) and A β 42 (right) were quantified as described above. The data were analyzed by one-way analysis of variance followed by Tukey's test (* $P < 0.05$; ** $P < 0.01$; ns, not significant).

In contrast, the Alc α ICD(AWAA) mutant failed to inhibit KLC1 association with JIP1b, did not attenuate the recovery of JIP1b and APP (Figure 6A), and showed no effect on APP transport (Figure 4E). Finally, overexpression of Alc α ICD(WT) in differentiating CAD cells dramatically inhibited anterograde transport of APP-GFP-containing vesicles and resulted in the aggregation of APP-containing vesicles (Figure 4D). These observations suggest that excess Alc α ICD attenuated the connection of APP to kinesin-1, causing a defect in APP transport.

Stokin *et al* (2005) suggested that deficits in axonal transport induce early pathogenesis in AD. Thus, we tested whether APP-containing vesicles actively transported by the kinesin-1 motor could suppress the generation of A β 40 and A β 42, compared to those that have stopped moving after dissociation from kinesin-1 in the presence of an excess amount of Alc α ICD. To analyze the change in A β generation more quantitatively, we used APP Swedish mutant (APPsw) (Citron *et al*, 1992), which is more susceptible to cleavage to produce greater amount of A β than the wild-type APP. The APPsw was transfected into HEK293 cells with or without JIP1b, KLC1, or Alc α ICD and the amount of secreted A β 40 and A β 42 was quantified by sELISA. Up to 10-fold greater amounts of A β 40 and A β 42 were generated after transfecting APPsw into cells, compared to the level of endogenously generated A β (Figure 6B and C). KLC1 expression did not affect the generation of A β 40 and A β 42, although JIP1b expression showed a slight decrease in generation of both A β species, as reported previously (Taru *et al*, 2002a). The levels of A β 40 and A β 42 secreted from APPsw cells decreased $\sim 25\%$ following expression of JIP1b and KLC1 (Figure 6C), which is consistent with the level of JIP1b expression alone (Figure 6B). This suppression was reversed by coexpression of Alc α ICD(WT) or Alc α ICD(NPAA); however, Alc α ICD(AWAA) coexpression did not block the suppressive effect of JIP1b and KLC1 on A β 40 and A β 42 generation. These observations suggest that JIP1b connection of APP to KLC and the resulting kinesin-1-dependent active movement of APP-containing vesicles suppress generation of A β 40 and A β 42, and that stationary and/or aggregating states of these vesicles increase A β 40 and A β 42 generation. These observations support the previous report by Stokin *et al* (2005) and suggest a significant role of JIP1b and Alc α in APP transport and A β generation.

Disruption of axonal transport of APP-like-containing vesicles by dAlc perturbs motor neuron activity

Coordinate sharing of Alc α cargo vesicles and APP-containing vesicles by kinesin-1 may be necessary for normal neuronal function. To analyze these functions, we expressed the *Drosophila* ortholog of alcadein (dAlc) in flies. UAS-dAlc was expressed in all *Drosophila* neurons using the panneuronal *elav-Gal4* driver (Chalvet and Samson, 2002). Third-instar larvae overexpressing dAlc showed locomotor defects, including aberrant tail-flipping movements, rather than normal waves of contraction (Figure 7, upper panels, and Supplementary Movie 7 in Sup_8.mov). These defects were not observed in *elav-GAL4/+* larvae not overexpressing dAlc. Immunostaining of larval segmental nerves, which connect to muscles, with an anti-synaptic vesicle-marker protein (CSP) and anti-*Drosophila* APP-like (APPL) antibodies revealed an accumulation of enlarged and aggregated vesicles in *elav-GAL4/UAS-dAlc* axons. These enlarged

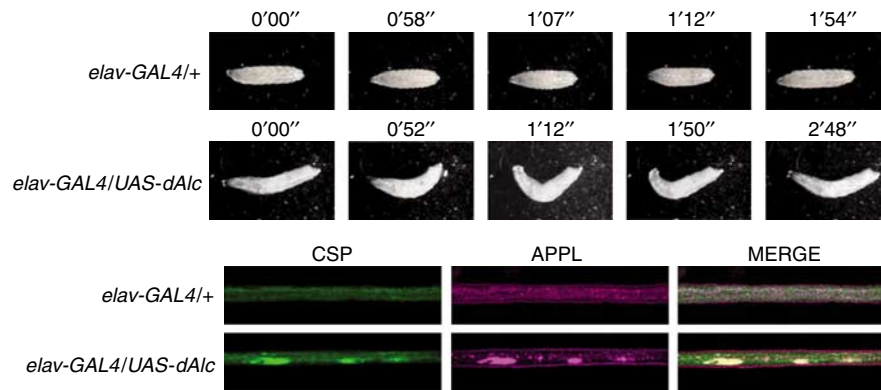


Figure 7 Impairment of motor neuron function and axonal transport by dAlc expression in *Drosophila*. (Upper panels) Larval movement. Movement of *elav-GAL4/+* (upper) and *elav-GAL4/UAS-dAlc* (lower) third instar larvae. Numbers indicate time. See Supplementary Movie 7, in Sup_8.mov. (Lower panels) Aggregation of APPL-containing vesicles by dAlc expression. Immunostaining of axons from *elav-GAL4/+* (upper) and *elav-GAL4/UAS-dAlc* (lower) third instar larvae. Larval segmental nerve axons were stained with antibodies against a synaptic vesicle marker protein (CSP) and APPL. Merged signals are shown in the panels on the right.

vesicles contained APPL (Figure 7, lower panels) and resemble APP-containing vesicles formed in differentiating CAD cells expressing APP-GFP in the presence of Alc α 1 and Alc α ICD (Figure 4C and D). The disruption of APPL anterograde transport by overexpression of dAlc may disrupt motor neuron function.

Discussion

Here, we reported five novel findings relevant to the function of Alc α and APP metabolism. (i) Alc α binds directly to KLC, and Alc α cargoes are transported by kinesin-1 at speeds slower than the transport velocity of APP-containing vesicles, which are also transported by kinesin-1. Thus, the velocity of transport is not solely determined by motor molecules but may be dependent on the combination of motor molecules and cargoes or adaptor proteins. (ii) Alc α can induce vesicle association of the kinesin-1 motor for axonal transport of cargo. The ability to regulate vesicle association of kinesin-1 motor components in the axonal transport of the cargoes is a novel feature of Alc α that has not been observed in other cargo and/or adaptor proteins. (iii) Alc α and JIP1b compete for kinesin-1 binding. (iv) Transport of APP-containing vesicles by kinesin-1 suppresses A β generation, while accumulation of stationary or aggregated APP-containing vesicles, potentially resulting from competition for KLC binding by another cargo, causes increased A β generation. (v) Disruption of the normal kinesin-1 motor system sharing by cargo or adaptor proteins produces defects in neural function. This is the first identification of Alc α as a novel cargo regulating kinesin-1 function and the first analysis of its association with APP transport and metabolism.

A schematic representation of the activity of Alc as a cargo is shown in Supplementary Figure S8A. Secretory vesicles containing Alc α activate kinesin-1 motor components through interactions with KLC and initiate anterograde transport of cargo. At the appropriate destination, KLC must release the Alc α cargo from the microtubule track and Alc α then appears on the plasma membrane. The slow dissociation of the KLC–Alc α interaction suggests that a covalent modification may be necessary for dissociation. In contrast, kinesin-1 transports APP-containing vesicles via JIP1b

(Supplementary Figure S8B). JIP1b may connect KLC to APP or to an unidentified cargo (Supplementary Figure S8B, left panel). Kinesin-1 may release the APP-containing vesicle, replacing it with an Alc cargo vesicle. Alternatively, Alc α ICD resulting from Alc α cleavage could compete with JIP1b or an unknown cargo for KLC binding, which would also release APP-containing vesicles (Supplementary Figure S8B, middle panel). Competition for kinesin-1 may promote accumulation of APP-containing vesicles in axons and cell bodies and facilitate A β generation (Supplementary Figure S8B, right panel). It is still unclear whether APP-containing vesicles include APP cleaving enzymes such as β -secretase and/or γ -secretase (Kamal *et al*, 2001; Lazarov *et al*, 2005). APP-containing vesicles aberrantly released from kinesin-1 may fuse more easily to other vesicles containing APP secretases in the aggregated state or be more easily transported to membrane regions rich in secretases.

It is still unclear whether APP serves as a cargo or not. Kamal *et al* (2000) reported a direct interaction of APP with KLC (Kamal *et al*, 2000), whereas our study and other groups have indicated that KLC most likely interacts with APP through JIP1b (Matsuda *et al*, 2003; Inomata *et al*, 2003). However, neither of these two conflicting observations prove or disprove the hypothesis that APP is a cargo of kinesin-1. However, a report that included the combined work of several research groups did not support the hypothesis that APP acts as a kinesin-1 cargo (Lazarov *et al*, 2005). Whatever the outcome of this polemic, our results clearly show that APP is transported by the kinesin-1 motor system and that this transport competes with Alc transport for kinesin-1.

In neurons, APP and Alc form a complex through cytoplasmic interactions with X11L, and both APP and Alc are metabolically stable in the complex (Araki *et al*, 2003, 2004). However, Alc α transport was largely independent of APP transport. Immunostaining of transporting vesicles in axon indicated that \sim 30% of vesicles contained both APP and Alc α and the other \sim 70% of vesicles contained either APP or Alc α . One possible interpretation is that APP and Alc are separately transported to nerve terminals, and that vesicles containing either protein are released at their destination and then exposed on the plasma membrane. APP, X11L, and Alc may form a complex either before leaving the Golgi appara-

tus, at the plasma membrane, or during the endocytotic process. Our preliminary immunostaining observations in differentiating CAD cells suggest that APP, X11L, and Alc α were largely colocalized in the Golgi, whereas APP and X11L did not show any apparent colocalization of vesicles transporting along the axon, although X11L was diffusely distributed in the cytoplasm with some accumulation around the Golgi (unpublished observation).

Prominent defects in axonal transport are often observed in neurodegenerative diseases. Disorders in axonal transport are often accompanied by axon terminal enlargement and aberrant accumulation of cargo (Yagishita, 1978). APP and Alc are highly colocalized in dystrophic neurites (Araki *et al*, 2003), which are enlarged by the accumulation of transported vesicles in the AD brain (Shoji *et al*, 1990; Cras *et al*, 1991). Moreover, in a fly model, overexpression of *Drosophila* APPL results in disruption of axonal transport, suggesting a role for APPL in microtubule-dependent vesicle trafficking (Torroja *et al*, 1999). Our data support the hypothesis that APP and Alc share the same kinesin-1 motor system for transport and that disruption of axonal transport of APP-containing vesicles can enhance the generation of A β , which may be a primary cause of AD pathogenesis (Stokin *et al*, 2005). Axonal transport well coordinated by the kinesin-1 motor for APP-containing vesicles and Alc cargo vesicles may be required for normal neuronal function as suggested by our observations that neuronal expression of dAlc in flies disrupts axonal transport of APPL-containing vesicles and perturbs motor neuron activity. These studies indicate that a quantitative imbalance in dAlc and APPL, and potentially APLIP1, which is a *Drosophila* JIP1b ortholog that associates with APPL and kinesin (Taru *et al*, 2002b; Horiuchi *et al*, 2005), results in vesicular aggregation and the accompanying deficits in neuronal function. Furthermore, dysfunctions in the exquisitely regulated axonal transport system may disturb formation of a stable tripartite APP, X11L, and Alc complex and result in overproduction of A β (Araki *et al*, 2003). These and future analyses of the molecular mechanisms of transport of APP-containing vesicles and Alc cargo vesicles should shed light on AD pathogenesis as they elucidate cargo-transport mechanisms.

Addendum

During the review process of this manuscript, similar observation on Calsyntenin/Alc binding to kinesin-1 was reported by Konecna *et al* (2006).

Materials and methods

cDNA cloning and plasmid construction

Mouse KLC1 cDNA encoding 542 amino acids (GenBank accession number AY753300/AK031309) and KLC2 cDNA encoding 619 amino acids (BC014845) were cloned into the pcDNA3.1 vector (Invitrogen) at the *Hind*III/*Xba*I sites for KLC1 (pcDNA3.1-mKLC1) and the *Eco*RI/*Eoc*RV sites for KLC2 (pcDNA3.1-mKLC2). The KLC1 used in this study has a 98% DNA sequence homology and a 97% protein sequence identity to KLC1A (BK000668). Further information is described in Supplementary data, Sup_1.pdf.

Antibodies

The anti-KLC1 polyclonal rabbit antibody (UT109) was raised against a peptide of mouse KLC1, amino acids 154–171, with an added amino-terminal Cys (C + KKYDDDISPSEDKDSDS), and the anti-KLC2 polyclonal rabbit antibody (UT110) was raised against a peptide of mouse KLC2, amino acids 149–163, with an added

N-terminal Cys (C + DASPNEEKGDVPKDS). Further information is described in Supplementary data, Sup_1.pdf.

Total internal reflectance fluorescence microscopy analysis

The cells were observed using a total internal reflectance fluorescence (TIRF) microscopy system (C1; Nikon) with a CCD camera (Cascade 650; Photometrics Co.). The transport velocity of vesicles was analyzed with Metamorph 6.1 software (Universal Imaging Co.), and the average values of 10 frames at 667 ms/frame were plotted. Further information is described in Supplementary data, Sup_1.pdf.

In vitro binding assay and SPR analysis

cDNA clones encoding the cytoplasmic domain of human Alc α (amino acids 873–971 of Alc α 1) and deletion constructs (Figure 1F, left panel) were subcloned into pGEX-4T-1 (GE Healthcare) to produce the GST-Alc α cyt fusion proteins. *In vitro* binding assays using GST fusion proteins were performed (Taru *et al*, 2002b) as described in Supplementary data, Sup_1.pdf. SPR analysis was performed using the Biacore system (Biacore International), as described in Supplementary data, Sup_1.pdf.

Co-immunoprecipitation and Western blot assays

Preparation of cell lysates and brain membrane fractions is described in Supplementary data in Sup_1.pdf. The lysates and their immunoprecipitates were subjected to Western blot analysis using specific antibodies. To accomplish more complete separation of the two Alc α bands, we used a discontinuous separating gel composed of 8% (w/v) and 15% (w/v) polyacrylamide with a standard stacking gel.

siRNA knock down of KLC1 and KLC2

cDNAs of siRNA were designed against the mouse KLC1 (nucleotides 1433–1451, 5'-GTTTGAAGCTGCAGAGACA-3') and KLC2 (nucleotides 1193–1201, 5'-GGAGTTTGGCTCAGTCAAT-3') genes and subcloned into the pSuper vector (OligoEngine). Mouse CAD cells and primary cultured neurons were cotransfected with GFP (pcDNA3.1-GFP) and the expression level of endogenous KLC1 and KLC2 was examined by immunostaining of transfected cells expressing GFP with the UT109 and UT110 antibodies.

Immunostaining of CAD cells and mouse peripheral nerves

CAD cells, a mouse CNS catecholaminergic cell line in which morphological differentiation can be initiated by serum depletion (Qi *et al*, 1997), expressing Alc α 1 and FLAG-KLC1 were fixed and stained with the UT83 and M2 antibodies. After the cells were washed, they were further incubated with an Alexa Fluor 488-conjugated goat anti-rabbit IgG or an Alexa Fluor 546-conjugated goat anti-mouse IgG. The cells were viewed using a confocal laser scanning microscope (LSM510 META NILO, Carl Zeiss). Immunostaining of mouse peripheral nerves is described in Supplementary data, Sup_1.pdf.

Drosophila study

cDNA encoding dAlc was purchased from Invitrogen (ResGen product) and inserted into pUAST (Brand and Perrimon, 1993) to construct pUAST-dAlc. Germline transformation was carried out as described previously (Robertson *et al*, 1988). The homozygous UAS-dAlc insertion line or Canton S (wild type) was crossed with the *elav*-GAL4 line and two independent P-inserted lines were analyzed, both showing identical phenotypes.

Supplementary data

Supplementary data are available at *The EMBO Journal* Online (<http://www.embojournal.org>).

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