Synaptic scaffolding protein SYD-2 clusters and activates kinesin-3 UNC-104 in C. elegans

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Kinesin-3 motor UNC-104/KIF1A is essential for transporting synaptic precursors to synapses. Although the mechanism of cargo binding is well understood, little is known how motor activity is regulated. We mapped functional interaction domains between SYD-2 and UNC-104 by using yeast 2-hybrid and pull-down assays and by using FRET/fluorescence lifetime imaging microscopy to image the binding of SYD-2 to UNC-104 in living Caenorhabditis elegans. We found that UNC-104 forms SYD-2-dependent axonal clusters (appearing during the transition from L2 to L3 larval stages), which behave in FRAP experiments as dynamic aggregates. High-resolution microscopy reveals that these clusters contain UNC-104 and synaptic precursors (synaptobrevin-1). Analysis of motor motility indicates bi-directional movement of UNC-104, whereas in syd-2 mutants, loss of SYD-2 binding reduces net anterograde movement and velocity (similar after deleting UNC-104’s liprin-binding domain), switching to retrograde transport characteristics when no role of SYD-2 on dynein and conventional kinesin UNC-116 motility was found. These data present a kinesin scaffolding protein that controls both motor clustering along axons and motor motility, resulting in reduced cargo transport efficiency upon loss of interaction.

The neuron is a highly polarized cell that possesses dendrites that are specialized for signal reception, and an axon for conduction and transmission. In axonal presynaptic termini, proper vesicle pool organization at the “active zone” and recruitment of synaptic vesicles apposing postsynaptic receptors is completed by complex interactions of presynaptic proteins, including SYD-2/liprin-α. The syd-2 gene encodes a member of the liprin family of proteins (i.e., “LAR-interacting proteins”) that assembles into protein scaffolds that localize presynaptic proteins and mediates targeting the presynaptic transmission machinery to opposite postsynaptic densities (1). It was reported that defects in the syd-2 gene cause a diffuse localization of synaptic vesicle markers in conjunction with a lengthening of presynaptic active zones in Caenorhabditis elegans GABAergic DD and VD motoneurons (2) whereas a mutation in the coiled-coil domain promotes synapse formation dependent on ELKS (3). SYD-2 seems to play a key role in recruiting presynaptic components acting downstream of the synaptic guidepost protein SYG-1 (4). In addition to a scaffolding function at the synapse, Drosophila liprin-α mutants display synaptic vesicle transport defects (5).

The long-range transport of vesicle cargo to synaptic sites requires molecular motor proteins of the kinesin superfamily. UNC-104/KIF1A, a member of the kinesin-3 family, is an essential neuron-specific, monomeric motor that transports synaptic vesicle precursors via a motor/lipid interaction involving the motor’s pleckstrin homology (PH) domain (6). Mutations in C. elegans UNC-104 impair the anterograde transport of synaptic vesicles from the soma to the synapse which results in uncoordinated, slow body movements of the worm (7). Shin et al. (8) reported a direct interaction of liprin-α with KIF1A in vitro, suggesting that liprin-α may function as a KIF1A receptor that links the motor to various liprin-α-associated proteins such as glutamate receptor-interacting protein and AMPA glutamate receptors (9, 10). As the function of KIF1A/liprin-α interaction remains unknown, we evaluated the underlying mechanisms of UNC-104/SYD-2 interaction in vitro and in vivo. As SYD-2 is thought to be a cargo of UNC-104 (11), we hypothesize that the scaffolding protein SYD-2 might coordinate motor organization on the synaptic vesicle membrane, which could regulate anterograde cargo transport (12, 13).

Results

The functional interaction between UNC-104 and SYD-2 was studied in worms expressing UNC-104 fused to the N terminus of a fluorescent protein [GFP or monomeric red fluorescent protein (mRFP); supporting information (SI) Fig. S1]. We used 2 syd-2 mutant alleles: a point mutation in glutamine 397 leading to a stop codon in the coiled-coil region (named ju37, ref. 2) and a deletion covering most of the N-terminal coiled-coils (named ok217). Note that the graph in Fig. 1 A shows averages of relative mRNA levels [based on real-time quantitative PCR (qPCR) experiments] and the gel (Fig. 1A Upper Left) shows a selected single RT-PCR experiment. Thus, band intensities in the gel do not necessarily reflect the average mRNA levels as we have determined by qPCR. Sequencing the ok217 allele revealed a missense mutation leading to an ochre stop codon at position 200. To test for expression of truncated SYD-2 products, we detected full-length proteins in N2 lysates and the corresponding protein fragment in syd-2(3ju37) by Western blotting [Fig. 1 A; UNC-104::GFP(ju37)]. However, no ok217 fragment (1–200 aa) was detected [lanes UNC-104::GFP(ok217) and ZM607(ok217)]; possibly because of degradation of this small protein trunk in the worm (even though high mRNA levels were detected; Fig. 1 A). Thus, the ok217 allele may represent a null allele, whereas ju37 shows detectable levels of both mRNA and truncated protein. Yeast 2-hybrid analysis was performed to test interaction domains of UNC-104 and SYD-2. Based on the known interaction of liprin-α and KIF1A (8), the motor’s stalk and liprin’s coiled-coil domains are prime candidates for in vitro binding testing. In addition to the previously published interactions, we found that either all (1–695) or some coiled-coils (341–695) of SYD-2 weakly interact with UNC-104 domain constructs (Fig. 1B and Fig. S1). The most prominent interaction occurs with the C-terminal half of SYD-2, including the SAM domains and UNC-104 stalk and FHA


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domain; still, interestingly, region 1–397 (corresponding to \textit{ju37} allele) also interacts with the FHA and stalk domains. However, the interaction of a nearly full-length SYD-2 (13–1,087) with UNC-104 is reduced. Direct binding was confirmed by pull-down experiments with recombinantly expressed proteins (Fig. S1B). Based on these findings, we assume that SYD-2’s coiled coils can intramolecularly interact with its SAM domains, thus masking potential UNC-104 binding sites (14). These data suggest that SYD-2 and UNC-104 can interact through multiple domains, while strong interactions occur between the SYD-2’s coiled coils 5–8 and SAM domains and UNC-104’s FHA and stalk domains, respectively.

**In Vivo FRET/Fluorescence Lifetime Imaging Microscopy Experiments Reveal a Close SYD-2/UNC-104 Interaction.** Fluorescence lifetime imaging microscopy (FLIM) measurements were used to determine whether SYD-2 and UNC-104 are able to physically interact in the living worm. Head neurons of the worm expressing both SNB-1::GFP and UNC-104::mRFP showed no FRET as the fluorophores are on opposite sides of the vesicle membrane. Values represent mean ± SEM. (Scale bar, 10 μm.)

Worms expressing only GFP::SYD-2 (Fig. 1E) exhibit a fluorescence lifetime of 2.74 ns ± 0.01 (\textit{n} = 11 worms), typical for GFP in the absence of FRET and significantly higher than in worms expressing both GFP::SYD-2 and UNC-104::mRFP (Fig. 1D; color bar and lifetime histograms in Fig. 1F; also see SI text). As a negative control, worms expressing both SNB-1::GFP and UNC-104::mRFP were analyzed by FRET/FLIM. Although motor and cargo vesicles co-localize under epi-fluorescence observation (Fig. 4G), the 2 respective fluorophores should not be in close proximity as the GFP of the synaptobrevin is located inside the vesicle membrane (15), whereas the mRFP of the UNC-104 is located outside (Fig. 1G). In summary, these results reveal an in vivo interaction of UNC-104 with SYD-2 in the living worm that appears more pronounced in younger animals.

**SYD-2 Regulates UNC-104 Motor Motility.** We then compared the transport characteristics of motors and vesicles in WT and \textit{syd-2} mutant backgrounds by UNC-104::GFP particle analysis in living worms in sub-lateral neurons (Fig. 2 C-F) and in isolated neurons (Fig. 3) by using spinning-disc confocal time-lapse fluorescence microscopy. Two examples of moving particles in a time series is shown in Fig. 2A (Movie S1) with corresponding positions indicated in the kymograph (Fig. 2B; 1-5, 6-9; x, static particle) and active anterograde and retrograde traffic events are shown in Fig. S3B. Most strikingly, in the living worm, anterograde velocity of (bidi-
vesicle-associated cargo SNB-1 undergoes a similar shift to retrograde transport parameters and overall velocity reduction in terminal truncated SYD-2; Fig. 2 phenotype in syd-2(ok217) mutants and switches from anterograde-based to retrograde-directionality in WT (Fig. 2A). In worms co-expressing UNC-104::mRFP5SYD-2::GFP, single UNC-104 particles (wt–coloc) have similar velocities than in ok217, but attain normal velocities when co-migrating with SYD-2 (wt–+syd-2 coloc). (F) Velocity, pausing, percentage of directionality, and net transport lengths are presented for UNC-104::mRFP in WT, syd-2(ju37), and syd-2(ok217) worms. (Scale bars, 10 sec for vertical/time, 10 μm for horizontal/distance.) Note that anterograde velocity of UNC-104 is reduced in SYD-2 mutants (ju37 and ok217), whereas retrograde velocity is increased in ok217. (F). For detailed discussion, refer to the text and SI text. Values represent mean ± SEM. *P < 0.05, **P < 0.01 (Student t test) comparing anterograde versus retrograde velocity. (Scale bar, 45 μm.)

From Fig. 2F it appears that UNC-104 activity is decreased in syd-2 mutants and switches from anterograde-based to retrograde-based movements. Indeed, motor particles exhibited increased net retrograde movements in syd-2 from a predominantly anterograde directionality in WT (Fig. 2F). Similarly, the total net transport (i.e., net displacement over one particle track) is decreased for anterograde events and increased in retrograde directions with a stronger phenotype in syd-2(ok217) (SYD-2 null) than in syd-2(ju37) (C-terminal truncated SYD-2; Fig. 2F).

UNC-104 in cell culture shows the same qualitative switch to retrograde transport parameters and overall velocity reduction in syd-2 mutants (Fig. 3 A-D). Intriguingly, UNC-104’s anterograde vesicle-associated cargo SNB-1 undergoes a similar shift to retrograde motility (Fig. 3). As the motor stalk domain interacts with SYD-2 (Fig. 1B), we tested this construct for movement characteristics. UNC-104ΔSTALK shows qualitatively similar changes as observed for UNC-104 in the syd-2(ok217) background with reduced velocity and increased pausing duration (Fig. 3 A and B).

However, no difference in the ratio of anterograde/retrograde movement was detected (Fig. 3C) while the total net transport was reduced by 40%. Deletion of the motor head surprisingly showed no dominant negative effect in movement when expressed in N2 WT animals; however, we cannot rule out that motorless UNC-104 does not interact with endogenous motor. As the UNC-104ΔSTALK construct still includes the FHA domain capable of SYD-2 interaction (Fig. 1B), a partial phenotype might explain the anterograde preference. As UNC-104 and vesicles tend to move in anterograde rather than in anterograde direction, with a tendency to overall retrograde events (Fig. 3). DLC-1 movement characteristics were not altered when expressed in ok217 background with the exception of a reduced net transport (Fig. 3). As expected, the vesicle cargo maker SNB-1 undergoes a similar shift to retrograde motility (Fig. 3). Analysis of conventional kinesins (kinesin-I, UNC-116::GFP) velocity, directionality (not shown), and expression pattern shows no difference between WT versus syd-2(ok217) allelic background (Fig. S5).
analyzed. Values represent mean ± SEM. *P < 0.05 (Student t test).

**SYD-2 Scaffolding Protein Clusters UNC-104 in the Ventral and Dorsal Sub-Lateral Neurons.** To test whether, in the living worm, an interaction between SYD-2 and UNC-104 is involved in the axonal localization of UNC-104, we crossed UNC-104::GFP-expressing worms into different syd-2 mutant worms following distribution pattern analysis. Fig. 4 A and A′ shows a representative example of UNC-104 distribution with occasional small punctae and large clusters in the ventral and dorsal sub-lateral neurons. This clustering is significantly reduced in SYD-2-knockout worms (ok217 allele; Fig. 4B), while clustering still occurs (although it is less pronounced) in syd-2(ju37) with shorter truncation products (Fig. 4C). Quantification of cluster properties in neurites reveals a significant decrease in cluster density and an increase in cluster shape elongation in syd-2 mutants (Fig. S2 H and I and Fig. STG).

To answer the question whether UNC-104::GFP cluster may resemble en passant synapses, we analyzed the distribution pattern of synapses visualized by the synaptic marker synaptobrevin-1, SNB-1::GFP. Fig. 4D shows a more regular (i.e., “pearl string”-like) distribution pattern of synapses compared with UNC-104 particles (Fig. 4A) whereas UNC-104 still co-localizes with SNB-1 in motor clusters (Fig. 4G and Fig. S4). However, in syd-2 mutant sub-lateral nervous system (Fig. 4E and F inset), synapses are arranged more irregularly (also see Fig. S7 D, F, and G) than in WT, and accumulations can be seen, whereas a diffuse and elongated synaptic morphology is consistent with previous observations from GABAergic DD motor neurons (2). As expected, deletion constructions of UNC-104 lacking SYD-2 interaction domains as STALK and FHA fail to cluster and properly localize in the axon (while deleting the motor and the PH domain did not affect axonal clustering; see Fig. S2). Last, clustering seemed to be dependent on the developmental stages of the worm (Fig. S2 F and G), whereas FRAP experiments (Fig. S6 A and B) and time-lapse imaging (Fig. S6C) reveal that clusters are highly dynamic structures.

**Discussion**

We wonder why UNC-104 and SYD-2 exhibit multiple interaction domains as revealed by our yeast 2-hybrid assays and used bioinformatics tools to reveal whether SYD-2 would belong to a class of proteins with “intrinsically disordered structures” (16). Examples of intrinsically unstructured proteins (IUPs) include tau/MAP2, SNAP-25, α-synuclein, and neurofilament-H. They have in common a lack of 3D structure in vivo, and their unfolded character enables various functional modes. Proteins of this remarkable class are able to bind to several partners in a structurally adaptive process. We used PrDOS (Protein Disorder prediction system; http://prdos.hgc.jp/cgi-bin/top.cgi) to investigate whether either UNC-104 or SYD-2 would belong to this class of proteins. Interestingly, analysis of UNC-104 did not reveal any relation to the IUP class; however, SYD-2 can be indeed considered an IUP. Thus, we believe that SYD-2’s multiple interactions site on UNC-104 would result in its multifarious functioning based on the lack of an ordered structure.

Shin et al. (8) reported that a 455–1,104 amino acid construct of liprin-α binds best to KIF1A at amino acid position 657–1,105 (therefore named the liprin-α binding domain). We confirm these in vitro interactions between similar constructs (UNC-104 655–1,105 and SYD-2 608–1,078; Fig. 1; note that homology between SYD-2 and liprin-α is of 67% similarity and 51% identity; see ref. 2). In addition, we found that a nearly full-length construct of SYD-2 (13–1,087) shows a weaker interaction compared with the shorter 608–1,078 (Fig. 1B and Fig. S1F), possibly based on an intramolecular head-to-tail folding mechanism (17). Most strikingly, the FHA domain of UNC-104 and the C-terminal half of SYD-2 contributes highly to the motor-scaffold interaction (Fig. 1B and Fig. S2). The FHA domain is supposed to be involved in regulating a monomer-to-dimer transition of UNC-104 as a result of its position between the 2 coiled-coil domains (18). In the living worm, UNC-104 and SYD-2 seemed to be adjacent enough for a physical interaction. For discussion on FLIM/FRET experiments and on UNC-104/SYD-2 interaction mechanisms, please refer to the SI text.

**UNC-104 and Synaptic Vesicle Motility in Neurons.** Although UNC-104 is thought to move unidirectionally, bidirectional motion of GFP-tagged UNC-104 was observed in vivo (19) (Figs. 2 and 3). Consequently, we differentiated between anterograde and retrograde events. In addition, we differentiated between single event velocities (i.e., no changes of directions) and velocities of runs with several events and directional changes, and found that single event velocities are significantly higher than those with several events. For example, the velocity of UNC-104::GFP(WT) particles with one moving event exhibit an average speed of only 1.12 μm/s ± 0.4 in WT cells [vs. ju37 (0.56 μm/s ± 0.24) or ok217 (0.62 μm/s ± 0.26)], faster than particles with several events and several directional...
is significantly reduced in syd-2 (Wagner et al. PNAS Early Edition). In C. elegans, the distribution pattern of UNC-104, SNB-1, and SYD-2 is shown in Fig. 4. Axonal distribution pattern of UNC-104, SNB-1, and SYD-2. (A and A1) UNC-104::GFP clustering in the ventral and dorsal sub-lateral neurons (arrow). Arrowhead indicates vulva. (Scale bars, 100 μm in A, 25 μm in A1) (B) Clustering is significantly reduced in syd-2(ok217) worms and fewer cluster are observed in the ju37 mutant (C). (D) Pearl string-like distribution pattern of synapses (SNB-1::GFP) differs from the distribution pattern of UNC-104 clusters in A. (E) In syd-2(ok217) worms, synapses are arranged more irregularly (Top Inset) compared with the WT worm (D) and tend to accumulate in the terminal endings (Bottom Inset). Dashed inset represents an example of a frequently observed diffuse accumulation of SNB-1. (F) In ju37 worms, synapses appear elongated (arrowhead indicates vulva; F and F' Inset with dashed line shows another example of SYD-2 distribution). (G) Co-localization of UNC-104::GFP (green) and SNB-1::mRFP (red) in sub-lateral neurons. Arrowheads indicate synapses. (Scale bar, 25 μm in B-G.)

Fig. 4. Axonal distribution pattern of UNC-104, SNB-1, and SYD-2. (A and A1) UNC-104::GFP clustering in the ventral and dorsal sub-lateral neurons (arrow). Arrowhead indicates vulva. (Scale bars, 100 μm in A, 25 μm in A1) (B) Clustering is significantly reduced in syd-2(ok217) worms and fewer cluster are observed in the ju37 mutant (C). (D) Pearl string-like distribution pattern of synapses (SNB-1::GFP) differs from the distribution pattern of UNC-104 clusters in A. (E) In syd-2(ok217) worms, synapses are arranged more irregularly (Top Inset) compared with the WT worm (D) and tend to accumulate in the terminal endings (Bottom Inset). Dashed inset represents an example of a frequently observed diffuse accumulation of SNB-1. (F) In ju37 worms, synapses appear elongated (arrowhead indicates vulva; F and F' Inset with dashed line shows another example of SYD-2 distribution). (G) Co-localization of UNC-104::GFP (green) and SNB-1::mRFP (red) in sub-lateral neurons. Arrowheads indicate synapses. (Scale bar, 25 μm in B-G.)

The speed of SNB-1/VAMP::mRFP transfected in C. elegans primary neuronal cell cultures (highest, 0.68 μm/s; lowest, 0.21 μm/s; average, 0.3 ± 0.13 μm/s; n = 327) was similar to the velocity of SNB-1/VAMP::GFP transfected in hippocampal neurons (up to 0.5 μm/s; compare ref. 21). DLC::YFP or SNB-1::mRFP expression in cells isolated from unc-104 mutants (e1265) did not reveal enough moving events for statistical useful studies. Furthermore, no directional motion was determined for a UNC-104ΔMOTOR::GFP construct transfected in cells isolated from unc-104 mutants (e1265). Thus, the transgene was crossed into a N2 (WT) worm, resulting in mobile UNC-104 with a deleted motor domain that is highly reduced even in the presence of endogenous fully functional motors (Fig. 3 Right). Probably, the truncated motor is still able to associate with endogenous motors, and detected moving characteristics may derive from a mixed motor population (i.e., functional and non-functional).

The finding that synaptic vesicles move slower than UNC-104 (SNB-1; Fig. 3A) favors a model wherein multiple motors are attached to a vesicle and are probably involved in a “tug of war.” Evidence for a functional dynein/UNC-104 cargo interaction was reported by Koushika et al. (22), showing that the transport of synaptobrevin, synaptotagmin, and UNC-104 requires the dynein heavy and light intermediate chain, respectively. In cultured neurons, DLC-1 velocity, pausing and directionality are unchanged but with reduced net transport in syd-2 mutants (Fig. 3), suggesting that dynein-based transport is only slightly affected by SYD-2. So far, a direct interaction between SYD-2/liprin-α and dynein could not be proven (5), but an indirect interaction via kinesin-I cannot be excluded. As the velocity of kinesin-I in syd-2 mutants is unchanged compared with WT (Fig. 5C), the rescue of UNC-104 velocity in the presence of SYD-2 (Fig. 2E), together with our in vitro binding studies (Fig. 1 and Fig. S1) suggest that SYD-2 directly enhances anterograde characteristics of UNC-104.

Model for UNC-104/SYD-2 Interaction. We propose a model in which UNC-104 binding to SYD-2 possibly enhances clustering of UNC-104 on the vesicle surface through multiple interaction domains. When UNC-104 is unbound from its cargo, it might form the largely observed diffuse transport domains (Fig. 3B). When UNC-104 is bound to SYD-2, it might aggregate into a N2 (WT) worm, resulting in mobile UNC-104 with a deleted motor domain that is highly reduced even in the presence of endogenous fully functional motors (Fig. 3 Right). Probably, the truncated motor is still able to associate with endogenous motors, and detected moving characteristics may derive from a mixed motor population (i.e., functional and non-functional).

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Materials and Methods

Constructs and C. elegans Strains. Generation of constructs and worm culturing were carried out according to standard protocols. We provide a thorough description of plasmid construction and the C. elegans strains used in the SI text.

Worm Lysates. Worm lysates were prepared from mixed-stage worms as described (23). In brief, three 6-cm plates were washed 3 times with M9 buffer and worms were resuspended in 100 mM ethanolamine, pH 8.0, 1 mM EDTA, including protease inhibitor mixture (Roche Diagnostics). Samples were boiled for 80 s and immediately resolved by 4% to 12% SDS/PAGE. Fifty micrograms of total protein lysates was loaded per lane. The polyclonal antibody against the N-terminal region 30 to 80 aa of SYD-2 was purchased from Santa Cruz Biotechnology.

RT-PCR and Real-Time qPCR. The primers of the 5' (forward, CAGAACGGAA-GATACCTGACTTCT; reverse, TGCCCAACAGGCCTCATT) end of the syd-2 gene covered a region upstream of the stop codon in the ok217 mutation (A2 bp). To evaluate the mRNA expression of the 3' end, we designed primers upstream of the SAM domains (1,802 bp; forward, CAACCAAGAAGCTTCAGGT; reverse, ACCTGGCGACCTGATGG). We took into account that, for RT-PCR experiments, primers need to cover at least one intron. As an internal control we designed primers covering the snb-1 gene. Real-time qPCR experiments were carried out based on the 2(-Delta Delta C(T)) method (24). We used the ribosomal protein rpl-18 gene as an endogenous control and N2 WT extracts as a “calibrator sample” (for details refer to ref. 24).

Bacterial Protein Expression and Purification. The fragment 623-1026 of UNC-104 was cloned using standard PCR methods into a pgEX-2T expression vector (Amersham/GE Healthcare). SYD-2 341–695 and 608-1089 fragments were expressed as fusion proteins to maltose binding protein (MBP) in a pMAL-2X expression vector (New England Biolabs). All constructs were verified by DNA sequencing. Proteins were expressed and purified by glutathione Sepharose chromatography (Amersham/GE Healthcare) or amylose resin (New England Biolabs) according to the manufacturer, followed by HiTrag-Q ion exchange chromatography (Amersham/GE Healthcare). Fusion proteins were either sayed or frozen with 10% sucrose added and stored in liquid nitrogen.

Yeast 2-Hybrid Assay. We used the Matchmaker GAL4 Two-Hybrid System 3 from Clontech (Invitrogen). Please refer to the SI text for detailed description of the yeast 2-hybrid assay analysis.

Primary Neuronal Cell Culture and Transfections. Primary cell culture was performed according to Christensen et al. (25). Primary neuronal cells with a cell density of approximately 65,000 cells per plate were transfected with either a pPD95.81::DLC-1::YFP or a pSM::Punc-86::SNB-1::mRFP construct by using TransFect transfection reagent (Promega) according to the manufacturer. Transfected wells were incubated at 22 °C in a humidified box for 1 to 2 d before microscopy.

Microscopic Transport Assay. Imaging was performed using a Zeiss Axiovert 200M microscope equipped with a QC100 spinning disk head and a Roper 512F EMCCD camera (Visitron). For a thorough description of our microscopic transport assay please refer to the SI text.

FRET/FLIM. Fluorescence lifetime sensing was performed by time-correlated single photon counting. The time-domain FLIM setup was used as an upgrade of a TSC-SP2 AOB5 laser scanning confocal microscope (Leica), equipped with a mode-locked femtosecond Ti:Sapphire Mira900 laser that is pumped by a Verdi-V laser (Coherent). The laser was tuned at 900 nm for 2-photon excitation of EGF (26). The fluorescence emission of EGF was detected using a band-pass filter centered at 515 nm ± 15 and placed in front of an MCP-PMT detector (R3809U-50; Hamamatsu Photonics). The acquisition board (SPC830) and software (SPCImage) were both from Becker & Hickl. Further analysis was performed by in-house-developed Matlab routines (MathWorks).

Statistical Analysis. Statistics of particle movement in the microscopic transport assay were carried out using the Student t test (two-tailed, unequal variance). Mean values are given with ± SEM if not marked otherwise. Statistical significance (confidence level) at a P value <0.05 is noted by asterisks.

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Supporting Information

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SI Materials and Methods

Constructs and C. elegans Strains. The UNC-104::GFP construct is a gift of Mimi Zhou and Jon Scholey (Davis, California) and has been described (1). C. elegans strains for analysis are derived from the unc-104(line CB1265 [unc-104(e1265) II]) and maintained at 20 °C to 25 °C using standard methods (2). Heritable lines of transgenic worms carrying extrachromosomal arrays of the constructs were created by microinjection of the Punc-104::UNC-104::GFP plasmid (70 μg/mL) into unc-104(e1265) hermaphrodites (3, 4). (Note the considerably high amount of this plasmid needed to barely rescue the unc-104 phenotype.) The UNC-104ΔMOTOR::GFP construct did not rescue the e1265 phenotype, and UNC-104ΔPH::GFP, UNC-104ΔFHA::GFP and UNC-104ΔSTALK::GFP constructs only mildly rescued the e1265 phenotype. For staining presynaptic vesicles, we used Punc-86::SNB-1::mRFP, whereas Posm-5::DLC-1::YFP was used as dynemin marker in cell culture transfections. The entire unc-116 gene without stop codon was cloned from genomic DNA into a Gateway expression vector Prab-3::GW-DEST (Invitrogen) and micro-injected with coinjection marker rol-6(su1006) into N2 and crossed into syd-2(ok217).

We crossed both UNC-104::GFP(e1265) and NM440(jsIs1) males into the following hermaphrodites carrying syd-2 mutations on the X chromosome: CZ9000/ii(su7) and ZM607(oki217) received from the Caenorhabditis Genome Center. Green fluorescent males from the F1-generation were back-crossed in syd-2 F0 hermaphrodites to obtain syd-2 homozygotes.

NM440(jsIs1) males were crossed into hermaphrodites expressing UNC-104::mRFP. The latter hermaphrodites were generated by micro-injection of the UNC-104::mRFP construct mentioned earlier in unc-104(e1265) worms, which rescued the uncoordinated phenotype. Males expressing pSM::Punc-86::GFP::SYD-2(wyk12) were crossed into UNC-104::mRFP-expressing hermaphrodites.

DNA Particle Bombardment. DNA micro-projectile bombardment was used to introduce pSM::Punc-86::SNB-1::mRFP and pPD95.77::UNC-104::GFP constructs at the same time into e1265 worms (rescuing the uncoordinated phenotype). Bombardments were performed using a PDS-1000/He unit from Bio-Rad and the protocol for DNA coating of gold particles and macro-carrier preparation described by Daines (5).

Yeast 2-Hybrid Analysis. Yeast transformation was performed according to the manufacturer’s protocol. Transformed cells were first plated on low-stringency selection medium (-LT) to identify double-transformants. Cells grown on –LT agar were either immediately replica-plated onto –HALT agar (high stringency) or first on –HLT-deficient media containing adnine (medium stringency) and then replica-plated onto –HALT agar (high stringency) plates. Cells that grow on selection media mostly tested positive for X-Gal. Those colonies that were negative for X-Gal but still grew on selection media received an overall lower score. The following scoring system was used: no colonies on –HALT or red or pinkish colony on –HALT, “-” (red color is a sign for adenine deficiency); white colony on –HALT, “+” or light blue colony on –HALT, “++”; and dark blue colony on –HALT, “+++.” For example, a –HALT plate (replica-plated from a –LT plate) with 1 white, 1 red, 3 light blue, and 4 dark blue colonies received an overall score of “+++” ((1 + 0 + (3*2) + (4*3))/9 = 2.1).

Microscopic Transport Assay and Image Analysis. Images were acquired at room temperature with a ×100 oil-immersion objective (N.A., 1.45) at 2 to 3 frames per second. Images were analyzed using ImageJ 1.36 software (National Institutes of Health; http://rsb.info.nih.gov/ij/). Kymograph images were obtained by drawing a line over the neurite of interest, followed by the application of the re-slice stack function. Static particles appear as vertical lines whereas the slope of moving particles corresponds to the velocity of the particle (Fig. 2A and B). The lines obtained for stationary particles (x in Fig. 2A and B) were used to correct for movement of the stage. Curved objects were straightened before their conversion to kymographs using the straighten curved objects tool from Kocsis et al. (6). Co-localization experiments were carried out using a DualView module (BFI Optilas). FRAP experiments (Fig. S6 A and B) were done on a Zeiss LSM510 Meta confocal microscope using the bleaching/time-lapse function.

For imaging, worms were immobilized by treatment with 5 mM levamisole (Sigma-Aldrich) before being placed on 2% agarose-coated objective slides.

Discussion

UNC-104/SYD-2 Interactions as Revealed by Yeast 2-Hybrid Assay. The intention of our yeast 2-hybrid assay was to provide complementary data to the in-depth yeast 2-hybrid analysis by Shin et al. (7). For example, we did not attempt to find minimal binding domains but to identify crucial domains for SYD-2/UNC-104 interaction.

First of all, we performed a thorough coiled-coil analysis using the coiled-coil prediction tool by Wolf et al. (8). We believe we provided a more accurate model of coiled-coil distribution for both UNC-104 and SYD-2 (Fig. S14). As the prey used by Shin et al. (7), KIF1A 455–1,105 aa, showed strong to very strong interaction, but the minimal binding domain KIF1A 657–1,105 aa showed a much weaker signal, we thought to investigate the function of a construct covering only the FHA domain (and the adjacent “second” coiled coil). Indeed, this construct interacts much more strongly with almost all SYD-2 constructs used in our study, especially compared with the “minimal binding domain” (LBD) KIF1A 655–1,105 aa. Further, we found that the minimal binding domain liprin-α 351–673 aa reported by Shin et al. (7) actually contained only a fragment of the last coiled coil (number 8 in our model). Thus, we designed a slightly longer fragment (SYD-2 341–395). Indeed, enlarging this fragment, interaction with almost all of our UNC-104 constructs was stronger (Fig. 1B). We also wanted to know whether the strong interaction of liprin-α 1–848 (and 1–673) is again based on coiled coil 8. Thus, we designed one construct covering cc8 (1–695), one that does not contain cc8 (13–608), and another one that also does not contain cc7. However, we could not identify any predominant role of cc7 or cc8 in these longer constructs. Last, we received a very different finding regarding the SYD-2 SAM domains. Shin et al. (7) showed a negative interaction between a liprin-α construct basically covering the SAM domains (688–1,202) and a KIF1A 455–1,105 construct. However, we report strong interactions of SAM domains with our UNC-104 constructs. Moreover, in a full-length SYD-2, all interactions are reduced in their strength, proposing that important SYD-2 domains might be masked as a result of an intramolecular SYD-2 interaction [as reported by Serra-Pages et al. (9)]. In summary, Y2H findings are as follows (Fig. 1B and Fig. S1): the FHA domain alone seemed to be a more important interaction partner than the
657–1,105 LBD; and SYD-2 coiled coil 8 and SAM domains (if not covered) are important domains for the interaction with UNC-104.

**FRET/FLIM Efficiency in Larvae versus Adult Animals.** Close inspection of FRET/FLIM data presented in Fig. 1G reveals that difference in FRET efficiency between younger and elder animals arises from the presence of variable amounts of individuals that exhibit no FRET, rather than from reduced overall FRET efficiencies. The positive cases in the adult group accounted for only one third of the total measurements (Fig. 1G), whereas only one animal in the larval group had negative findings. In those groups, the FRET efficiencies of positive animals were 6.0% ± 1.2% (n = 4 worms) and 7.5% ± 1.4% (n = 9 worms), respectively. These differences are not statistically significant.

We hypothesize that FRET efficiency is a function of SYD-2 phosphorylation: as SYD-2 undergoes an intramolecular folding if unphosphorylated (9), we assume that UNC-104 binding sites might be masked. Conversely, phosphorylated SYD-2 would be increased in larvae compared with adults (L1-L3, 4–5 copies; L4-adults, 1–2 copies), consistent with our assumption. However, it is still unclear why some adult animals show normal FRET efficiencies.

**Comparison of Kinesin-3 Movement in Neurons.** Lee et al. (12) reported KIF1A movements in rat hippocampal neurons of approximately 0.1 µm/s higher than shown here, whereas we determined average durations of persistent anterograde and retrograde movements were lower compared with those in the study of Lee et al. (12) (see Table S1). How can these differences be explained? We found that, for every experimental condition, there is the need to determine individual pausing characteristics of the observed particle, which might vary in size and shape. Analyzing our data, we determine a "pause" as a movement lower than (on average) 0.065 µm/s (0.05–0.08 µm/s). We assume that variations in published velocities for KIF1A/UNC-104 might be based on differences in determining this threshold. Also, for UNC-104 velocities in the living worm, we determined lower values than others. Considering the moving speed resulting from particles that did not make any directional changes, the average speed was 0.89 µm/s ± 0.43, which is also lower than that reported by Zhou et al. (1) (1.02 µm/s; velocities include directional changes). However, Zhou et al. (1) considered velocities lower than 0.2 µm/s as pauses, which were excluded from calculations. Moreover, a closer look at the kymographs published by Zhou et al. (1) (and the quotient of numbers of events and total numbers of measured particles) reveals that mostly single-event moving particles were probably analyzed (1.78 events per particle with 1,634 events total and 917 particles total, compared with 6.95 events per particle with 3,196 events total and 460 particles total in the present study). The average pausing duration determined by Zhou et al. (1) was almost twice as low as we determined, which is consistent with the higher speed of UNC-104 determined by Zhou et al. (1) and the lower one we determined.

In preliminary experiments, we did not see any differences of UNC-104::GFP moving characteristics in axons compared with dendrites (data not shown); thus, we focused on axons only. Similarly, Zhou et al. (1) determined only small differences between axons (1.01 µm/s ± 0.53; n = 464), dendrites (n = 1.19 µm/s ± 0.38; n = 237), and axonal commissures (1.03 µm/s ± 0.37; n = 33). In addition, in dendrites, a switching between anterograde and retrograde movements might be simply based on the mixed orientation of microtubules (13).

Fig. S1. (A) Constructs: a schematic drawing of UNC-104::GFP motor is shown with deletion constructs covering the motor domain (ΔMOTOR) aa 1–356, the forkhead domain (ΔFHA) aa 463–592, the SYD-2/liprin-binding domain (ΔSTALK) aa 654–1,339 and the PH domain (ΔPH) aa 1,460–1,584. Coiled-coil regions are marked with a white box. SYD-2 protein is shown with coiled-coil regions (for prediction method, see ref. 8), SAM domains, and a C-terminal PDZ binding domain. ju37 mutation leads to a stop codon at position 397 after the first 7 coiled-coil domains, whereas we found that ok217 creates an additional Tyr-199 and a stop codon at position 200. (B) In vitro pull-down assays of GST::UNC-104 (623–1,026) with MBP, MBP::SYD-2 (341–695), and MBP::SYD-2 (608–1,089). (Top) Coomassie-stained SDS/PAGE of pull-down assays. (Bottom) Corresponding Western blot detection of UNC-104 by anti-GST antibody staining. To confirm the strong interactions between the 2 SYD-2 constructs 341–695/608–1,089 and UNC-104, we performed a pull-down experiment with recombinantly expressed and affinity-purified GST::UNC-104 (623–1,026) and SYD-2 (341–695 or 608–1,089) fused to MBP (Fig. S1B). As expected, GST::UNC-104 is pulled down by both MBP::SYD-2 341–695 and 608–1,089 but not MBP alone. During the experiments, we found that the SYD-2 sequence was updated in the worm database resulting in our SYD-2 608–1,089 construct. As both C-terminal constructs behave similarly in interaction strength, we however believe that 2 additional amino acids at the C terminus would not affect the overall interaction. (C–E) Examples of yeast growth on highly selective HALT plates. (Note that here we show examples of plates, whereas in Fig. 1B we summarize mean scores of 2–5 independently carried-out experiments.) (C) Representative interactions between the FHA domain of UNC-104 and SYD-2 deletions. (D) Representative plates from screens with the UNC-104 STALK domain (655–1,105) and SYD-2 deletions, as well as positive controls p53 and Large T. (E) SYD-2 self interactions including positive controls. In Fig. S1C we excluded the syd-2 1–397 construct as the growth is comparable to that in plate B. Similarly, we do not show syd-2 608–1097 as growth is comparable to that in A. (F) Yeast 2-hybrid results table with additional constructs.
Fig. S2. Distribution of UNC-104 deletion constructs in WT and syd-2 mutant worms. To find out whether specific UNC-104 domains are involved in SYD-2 depended clustering, we examined the distribution of UNC-104 deletions constructs lacking SYD-2 binding sites in WT and syd-2 worms. Compared with the UNC-104::GFP WT motor (A), a worm expressing UNC-104::STALK::GFP lacking a SYD-2 binding site (stalk domain 655–1,339) (B) shows no clustering in the sub-laterals (arrows) but strong accumulation of UNC-104 in the terminal endings. A similar effect was apparent in worms expressing UNC-104::GFP with a deleted FHA domain (C). In both cases, expression of the same constructs in the ok217 mutant worm did not reveal a different localization pattern (data not shown). UNC-104::PH::GFP is retained in the cell bodies (arrows) but also clusters in neurites (D). Clustering was strongly decreased in the ok217 background. Similarly, UNC-104::GFP with a deleted motor domain exhibits enhanced retaining of UNC-104 in the cell bodies and clustering (E), which was mildly reduced in the ok217 background. These results suggest that SYD-2 interacts with the UNC-104 stalk and FHA domain rather than with the UNC-104 PH or motor domain, consistent with our yeast 2-hybrid results (Fig. 1B and Fig. S1). (Scale bar, 10 μm.) Analysis of cluster development in C. elegans larval stages: (F) developmental emerging of UNC-104 clusters in ventral and dorsal sub-lateral neurons during the L3 larval stage of C. elegans. (G) In the syd-2(ok217) mutant, no clustering occurs in neither of the 3 larval states. (Scale bar, 10 μm.) (H–I) To address the question how the endogenous motor is distributed in neuronal processes, we performed immunofluorescence stainings. Endogenous UNC-104 (H) shows a uniform, cytosolic distribution with cluster-like accumulations (similar to UNC-104::GFP) that are less pronounced in syd-2 mutants (I) (see Fig. S7G). (Scale bar, 10 μm.)
Fig. S3. Analysis of co-migration of UNC-104 motor with SYD-2. (A) Co-localization and movement of UNC-104 and UNC-104/SYD-2 at the same time displayed in a kymograph. Arrowhead indicates static SYD-2 (green); short, thick arrow indicates static UNC-104 alone (red); long, thin arrow indicates yellow tracks (moving). The y axis is 300 s, x axis is 61 μm. (B) Top: Confocal section of a sub-lateral neuron. Bottom: Kymograph along the top neuron length showing stationary (vertical lines) and moving particles (tilted lines). Movements to the right are in anterograde direction. From kymograph data, transport parameters were calculated from moving particles from in vivo whole-worm acquisitions. (C) Summary table of data analysis.

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<th>Green (SYD-2)</th>
<th>Yellow (SYD-2/UNC-104)</th>
<th>Red (UNC-104)</th>
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<tr>
<td>Moving particles</td>
<td>0% (0/1)</td>
<td>62% (13/21)</td>
<td>67% (2/3)</td>
</tr>
<tr>
<td>Static particles</td>
<td>100% (1/1)</td>
<td>38% (8/21)</td>
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Fig. S4. Cryo-EM revealed co-localization of UNC-104 and vesicle marker SNB-1. To understand the nature and structure of the observed clusters, we examined immunogold-labeled UNC-104 and UNC-104 co-labeled with SNB-1 in ultra-thin cryosections by transmission EM. UNC-104::GFP was labeled with anti-GFP and protein A-gold particles (10 nm) (A–C), synaptobrevin with anti-SNB-1 and protein A-gold particles (5 nm) (D and E). (A–C) Motors cluster to membranous structures that are distinct from mitochondria (M) (A) and co-localize with spherical SNB-1-positives vesicles (B and C). (B, Bottom) vesicle structures co-localizing with UNC-104 resemble vesicles in transit. (Scale bar, 500 nm.) (E) Higher magnification of cryo-EM sections labeled with an anti-SNB-1 antibody (5 nm gold) and anti-GFP (UNC-104, 10 nm gold) showing co-localization of the 2 labels to single vesicles. These experiments show that UNC-104 motors indeed cluster to membranous, vesicular structures with a diameter of 40–50 nm; see table (E).
Fig. S5. Conventional kinesin-I localization and velocity is not affected by SYD-2. UNC-116::GFP (conventional kinesin heavy chain) expressed under the UNC-104 promotor shows diffuse, cytosolic staining with occasional clusters in WT axons (A). The UNC-116::GFP staining appears unchanged in syd-2(ok217) mutants (B). (C) Quantification of UNC-116::GFP velocity in vivo (living worm) show no significant difference comparing anterograde (black bars) and retrograde (white bars) movement in WT and syd-2(ok217).
Analysis of motor cluster dynamics. To answer the question if UNC-104 cluster might represent a simple accumulation of inactivated motors, we photo-bleached clusters and followed their fluorescence recovery over time. (A) Photo-bleaching of a cluster (arrow) and following the fluorescence recovery over 60 s time. (B) Average fluorescence intensities of 4 individual bleached cluster (error bars represent SD). Fluorescence intensities were normalized to pre-bleach cluster intensity. (Scale bar, 10 μm.) Experiments in A and B demonstrate a 50% to 60% mobile pool of unbleached and active motors newly accumulating into the cluster. These results provide evidence that UNC-104 cluster are dynamic membranous structures to which motors can accumulate (possibly serving as an active motor pool) and eventually move along the neurite. Interestingly, clusters are not solely stationary motor deposits, as we also occasionally observe whole clusters moving at velocities ranging from 0.1 to 0.3 μm/s during time-lapse imaging (C, arrows), similar to large UNC-104 particles (Fig. 2E and Fig. S3 A and C) with long pause durations. Note that in C individual moving clusters are highlighted in different colors and that an asterisk marks a stationary cluster. (Scale bar, 10 μm.)
Fig. S7. Line scan of axonal clusters and cluster statistics. Axonal clusters highlighted in the magnified inset boxes in Fig. 4 (Ai, B, C, D, E Bottom, F, and F’) have been analyzed by line scans (ImageJ, version 1.42q), intensity values have been background-subtracted, and net values plotted against the line distance. (G) Quantification of UNC-104 cluster distribution. Quantification of UNC-104 cluster distribution, size, lengthening (i.e., Feret diameter) and circularity. The total neurite length is the sum of all segments measured shown as mean ± SD. (*t test, e1265 vs. ok217; **t test, ok217 vs. ju37; ‡ t test, e1265 vs. ju37; n.s., not significant.)
Fig. S8. Model of SYD-2 stimulated UNC-104 transport. In WT worms, the anterograde transport of synaptic cargo (black arrows) is stimulated by SYD-2 binding to UNC-104. SYD-2 binding causes motors to cluster along the axon that most likely represent cargo-detached motors. Through multiple interaction regions and self-dimerization, SYD-2 can concentrate motors locally and increase persistent and fast movement. Truncation mutants of SYD-2 [syd-2(ju37)] partially retain their ability to bind with lower strength to UNC-104, thus resulting in a less pronounced anterograde flux (black arrows) with an increase in retrograde transport events (white arrows). With reduced affinity to UNC-104 and scaffolding sites, fewer motors are found in clusters. In the absence of SYD-2, transport is unbalanced toward retrograde events with a decreased flux.
**Movie S1.** Example of a time-lapse imaging sequence of UNC-104::GFP particles in a neurite of a living worm.

[Movie S1 (MOV)]
Table S1. Comparison of reported KIF1A/UNC-104 motility

<table>
<thead>
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<th></th>
<th>This study</th>
<th>Zhou et al. (2001)</th>
<th>Lee et al. (2003)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Anterograde</td>
<td>Retrograde</td>
<td>Anterograde</td>
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<tr>
<td>Velocity in vitro, μm/s</td>
<td>0.78 ± 0.35</td>
<td>0.44 ± 0.13</td>
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<td>Velocity with no directional change, μm/s</td>
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<td>Persistence of movement in vitro, s</td>
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<td>11.83 ± 5.36</td>
<td>5.35 ± 4.60</td>
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<td>Pause duration, s</td>
<td>8.89 ± 4.28</td>
<td>12.67 ± 4.05</td>
<td>6.2 ± 7.71 (average)</td>
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<tr>
<td>Velocity (in cell culture), μm/s</td>
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<td>0.59 ± 0.25</td>
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<td>Persistence of movement (in cell culture), s</td>
<td>7.34 ± 3.86</td>
<td>5.57 ± 1.9</td>
<td>NA</td>
</tr>
</tbody>
</table>

KIF1A/UNC-104 movement in the living worm, cell culture and species others from *C. elegans*. Mean ± SD. NA, not applicable; ND, not determined.