Molecular motor function in axonal transport in vivo probed by genetic and computational analysis in Drosophila

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ABSTRACT Bidirectional axonal transport driven by kinesin and dynein along microtubules is critical to neuronal viability and function. To evaluate axonal transport mechanisms, we developed a high-resolution imaging system to track the movement of amyloid precursor protein (APP) vesicles in Drosophila segmental nerve axons. Computational analyses of a large number of moving vesicles in defined genetic backgrounds with partial reduction or overexpression of motor proteins enabled us to test with high precision existing and new models of motor activity and coordination in vivo. We discovered several previously unknown features of vesicle movement, including a surprising dependence of anterograde APP vesicle movement velocity on the amount of kinesin-1. This finding is largely incompatible with the biophysical properties of kinesin-1 derived from in vitro analyses. Our data also suggest kinesin-1 and cytoplasmic dynein motors assemble in stable mixtures on APP vesicles and their direction and velocity are controlled at least in part by dynein intermediate chain.

INTRODUCTION A hallmark of the neuron is its polarized axon, which can extend for more than 1 m in humans. Within the axon, a wide variety of cargoes essential for the viability and function of the neuron must be transported along microtubules between the neuronal cell body and synapses (Grafstein and Forman, 1980). Understanding how molecular motor proteins drive this axonal transport is important to the understanding of a wide range of neurological diseases (Goldstein, 2003; Stokin and Goldstein, 2006; Chevalier-Larsen and Holzbaur, 2006; De Vos et al., 2008).

Axonal transport toward synapses (anterograde transport) is driven by microtubule plus end–directed kinesins, especially those from the kinesin-1–4 and 13 families (Goldstein and Yang, 2000; Lawrence et al., 2004; Hirokawa and Takemura, 2005). Transport toward the neuronal cell body (retrograde transport) is driven by microtubule minus end–directed cytoplasmic dynein (King, 2000; Vallee et al., 2004). Much information about the biochemical and mechanical mechanisms of kinesin-1 and cytoplasmic dynein movement has emerged from in vitro studies. For example, single
RESULTS

Establishing the properties of axonal APP vesicle movement in vivo

We probed the transport behavior of individual APP vesicles in vivo by using SG26.1 Gal4 (Gal4 gene enhancer trap strain) to drive the expression of UAS-APPYFP in Drosophila. Because the expression of SG26.1 Gal4 is confined to a subset of ventral ganglion neurons, APPYFP movement could be tracked with single-axon resolution within segmental nerves. Analysis of green fluorescent protein–tagged EB1 (EB1-GFP) transport in these neurons confirmed that movement away from cell bodies is microtubule plus end–directed and anterograde (Supplemental Figure S1A), consistent with a previous report (Stone et al., 2008). Using high spatial (0.126 micron) and temporal (10 Hz) resolution imaging, we characterized the movement of APP in an otherwise wild-type background (Figures 1, A–E, and S1, A–F, and Supplemental Movie S1; see Materials and Methods): 32.3% of the vesicles moved anterogradely and 18.2% moved retrogradely (Figure 1, A–C), whereas 40.4% were stationary and 9.1% reversed direction with an average switch frequency of 6.09 switches/min (Figures 1C and S1B). Switches generally did not involve pausing, consistent with a previous report of lipid droplet movement in Drosophila embryos (Welte et al., 1998). Thus directional switches are mediated by an apparently instantaneous change in the balance of active motors.

Anterograde velocities of APP vesicles depend on the amount of kinesin-1

Considerable evidence demonstrates that APP movement is driven by kinesin-1 (Koo et al., 1990; Ferreira et al., 1992; Amarutunga et al., 1993; Simons et al., 1995; Tienari et al., 1996; Kamal et al., 2000; Szodorai et al., 2009). We observed that a substantial fraction of APPYFP vesicles traveled at velocities faster than those observed for kinesin-1 motors in vitro, up to a maximum of 2.85 μm/s (Figure 1D), consistent with previous reports (Kaether et al., 2000; Stamer et al., 2002; Goldsberry et al., 2006).

Kinesin-1–driven movement in vivo, which is substantially faster than in vitro, may be related to a different regulation of the ATPase cycle in vivo. Alternatively, multiple kinesin motors may coordinate to generate faster velocities, as speculated by Kural and colleagues (Kural et al., 2005). The latter would be incompatible with the independence of velocity and motor number for kinesin-1 seen in vitro (Howard et al., 1989; Block et al., 1990; Spudich, 1990). Additionally, it would be inconsistent with more recent in vivo data of lipid droplet movement in Drosophila embryos, which suggest that neither velocity nor run length changes significantly with varying amounts of kinesin-1 (Shubeita et al., 2008).

To test the hypothesis that the velocities and run lengths of APP vesicles may be determined by the amount of kinesin-1, we performed genetic, biochemical, and statistical analyses to correlate the number of active kinesin-1 molecules with vesicle behavior. We tested four predictions made by this hypothesis. A first prediction is that the anterograde velocity distribution should be nonnormal with multiple modal peaks reflecting differential occupancy of active motors. We tested this possibility by statistical mode analysis (Fraley and Raftery, 2002) and found that the distribution was best fit by three modes (Figure 1E; see Materials and Methods). Thus the first prediction is fulfilled. Interestingly, multimodal cargo velocity distributions have been reported previously in motility assays under substantial mechanical load of viscoelastic drag and in intracellular transport of PC12 cells and Xenopus melanophores (Hill et al., 2004; Levi et al., 2006; Gagliano et al., 2010).

A second prediction is that increased velocities should correlate with increased run lengths (or reduced pause frequency, Spudich, 1990). Because our relatively short observation window of 15 s leads to truncation of run length for many vesicles (Figure S1C), we tested for a correlation of segmental velocity with pause frequency (see Supplemental Material). For controls, the anterograde correlation coefficient was −0.45, indicating that the faster the cargo, the lower
sign, a fraction at the interface of the 8/35 sucrose step consists of floated membranes and vesicles. We found that 50% khc gene reduction leads to ∼50% reduction in both KHC and KLC proteins in the 8/35 fraction (Figure 2, D and E), indicating that khc reduction leads to less kinesin-1 associated with vesicles. Under these conditions, we observed substantial decreases in anterograde APPYFP velocities (Figures 2F and S2A). Although the distributions were best fit by three modes (as in control), there were significant shifts in the relative contribution of each mode in kinesin-1 reductions (Figures 2, G and H, and S5B and Supplemental Table S1). Moreover, either khc or klc reduction led to significantly shorter run lengths, increased pause frequency, increased pause duration, and a larger fraction of stationary vesicles (Figure S2, B–H). Together these biochemical and genetic findings are consistent with the notion that the number of functional kinesin-1 motors assembled per cargo is reduced with kinesin-1 reduction, causing a decrease in the anterograde velocity and a shift in the occupancy of modal velocities (unlike lipid droplet transport in Drosophila embryos [Shubeita et al., 2008]).

A fourth prediction of the hypothesis that the velocities of APP vesicles may be determined by the amount of kinesin-1 is that the correlation of anterograde velocity and pause frequency should be preserved in kinesin-1 reduction. Indeed, the magnitude of the average probability of a pause. Thus the second prediction is also fulfilled.

A third prediction is that reducing the amount of kinesin-1 on vesicles should shift the distribution of velocities to lower modes without changing the mode centers. We tested this possibility by partially reducing kinesin-1 level, thereby circumventing the issues of organismal lethality and cellular defects reported in previous in vivo studies (Gindhart et al., 1998; Hurd and Saxton, 1996; Boylan and Hays, 2002). We used two null point mutation alleles for kinesin heavy chain (KHC), khc8 and khc20 (Saxton et al., 1991; Brendza et al., 2000), and one deletion allele for kinesin light chain (KLC), klc8ex94 (Gindhart et al., 1998). Western blot analysis of kinesin-1 heterozygotes (see Materials and Methods) confirmed that the removal of one functional khc or klc gene caused ∼50% reduction in both KHC and KLC proteins (Figure 2, A–C). Interestingly, khc reduction also resulted in KLC protein reduction, whereas klc reduction did not affect KHC protein levels. Thus KLC protein levels appear to depend on KHC but not vice-versa, consistent with previous work in Drosophila S2 cells (Ligon et al., 2004).

To test whether kinesin-1 reduction translates to less motor protein assembly on vesicles, we used bottom-loaded sucrose flotation step gradients (see Materials and Methods). In our experimental design, a fraction at the interface of the 8/35 sucrose step consists of floated membranes and vesicles. We found that 50% khc gene reduction leads to ∼50% reduction in both KHC and KLC proteins in the 8/35 fraction (Figure 2, D and E), indicating that khc reduction leads to less kinesin-1 associated with vesicles. Under these conditions, we observed substantial decreases in anterograde APPYFP velocities (Figures 2F and S2A). Although the distributions were best fit by three modes (as in control), there were significant shifts in the relative contribution of each mode in kinesin-1 reductions (Figures 2, G and H, and S5B and Supplemental Table S1). Moreover, either khc or klc reduction led to significantly shorter run lengths, increased pause frequency, increased pause duration, and a larger fraction of stationary vesicles (Figure S2, B–H). Together these biochemical and genetic findings are consistent with the notion that the number of functional kinesin-1 motors assembled per cargo is reduced with kinesin-1 reduction, causing a decrease in the anterograde velocity and a shift in the occupancy of modal velocities (unlike lipid droplet transport in Drosophila embryos [Shubeita et al., 2008]).

A fourth prediction of the hypothesis that the velocities of APP vesicles may be determined by the amount of kinesin-1 is that the correlation of anterograde velocity and pause frequency should be preserved in kinesin-1 reduction. Indeed, the magnitude of the
such as a tail-flip phenotype or animal inviability. Second, while axonal swellings were present in our kinesin-1 reduction genotypes, they were generally found in distal axonal regions, rather than in the more proximal region we analyzed. Third, nonselective poisoning would likely shift modal velocity values, whereas we observed fractional shifts in mode. Fourth, both kinesin-1–dependent and kinesin-1–independent transport would be expected to show impairment. To test the latter possibility, we analyzed the movement of synaptotagmin (SYT)—a kinesin-3 cargo (Yonekawa et al., 1998; Pack-Chung et al., 2007) tagged with GFP (SYTGFP), in khc<sup>+/+</sup> and klc<sup>8ex94/+</sup> animals. As expected, we observed that kinesin-1 reduction
correlation coefficients for all kinesin-1 reduction genotypes was in the range of −0.45 to −0.55, which is statistically highly significant (Figure 2I; see Supplemental Material). Thus all four predictions are fulfilled.

A potential concern is whether kinesin-1 reduction causes nonspecific global toxicity to axonal transport, including swellings, “organelle jams,” and blockage in transport (Hurd and Saxton, 1996; Martin et al., 1999; Gunawardena and Goldstein, 2001; however, see Pilling et al., 2006). Several observations suggest nonspecific global toxicity is an unlikely explanation for our data. First, there was no organismal phenotype characteristic of axonal transport defects, such as a tail-flip phenotype or animal inviability. Second, while axonal swellings were present in our kinesin-1 reduction genotypes, they were generally found in distal axonal regions, rather than in the more proximal region we analyzed. Third, nonselective poisoning would likely shift modal velocity values, whereas we observed fractional shifts in mode. Fourth, both kinesin-1–dependent and kinesin-1–independent transport would be expected to show impairment. To test the latter possibility, we analyzed the movement of synaptotagmin (SYT)—a kinesin-3 cargo (Yonekawa et al., 1998; Pack-Chung et al., 2007) tagged with GFP (SYTGFP), in khc<sup>+/+</sup> and klc<sup>8ex94/+</sup> animals. As expected, we observed that kinesin-1 reduction

![Image of a page from a scientific journal containing text and graphs related to molecular motor function.]
does not significantly change the total number of SYTGFP moving vesicles (see Figure 4I later in the paper) or the population percentages (Figure S6, G–J; see also Table S2). Furthermore, kinesin-1 reduction has no significant effect on retrograde velocity (Figure S6, B and D) or anterograde run length (Figure S6E). Although we observed a mildly significant decrease in anterograde segmental velocity for khc<sup>8ex94</sup>/+, this was not seen in klc<sup>8ex94</sup>/+ (Figure S6, A and C). Interestingly, a significant increase in retrograde run length was noted (Figure S6F), which may relate to the overall decrease in kinesin-1 related transport. Taken together, the SYTGFP data suggest that nonspecific global toxicity is an unlikely mechanism underlying the effects of kinesin-1 reduction on APP movement.

**Kinesin-1–driven anterograde velocity modes are relatively stable**

Current models of vesicle movement make different predictions about the stability of movement behavior during axonal transport. For example, models in which motor proteins actively associate and dissociate with vesicles to determine directionality predict that certain parameters will vary over time (e.g., before and after pauses). Alternatively, models that invoke stable loading of vesicles in neuronal cell bodies predict substantial stability of movement before or after pauses and during runs. In fact, a prediction of the hypothesis that varying kinesin-1 motor amounts on APP vesicles control their behavior is that while the distribution of velocities is broad, the velocity of any given vesicle should remain relatively stable if motor number does not change over time. Hence, we analyzed movement behaviors relative to pauses and run lengths.

Our measurements of pause frequency indicate that APP vesicles pause infrequently during movement (Figure S1, D–E). Consistent with this, the observed average run length for APPYFP vesicles was 7.63 μm (Figure S2B), severalfold higher than in vitro (Howard et al., 1989; Block et al., 1990). Because our cargo trajectories are truncated spatially and temporally, even this value is too low. Our calculated estimated bulk anterograde run length is 11.7 μm (see Supplemental Material). Indeed, longer-duration movies (2 min of continuous imaging) generated an average run length of 21.85 μm (SEM = 2.3 μm; n = 47 segments; unpublished data).

We also probed the stability of APP movement by calculating the probability that a vesicle changes velocity mode during runs and after pauses and comparing it with random changes (see Supplemental Material). Consistent with the idea of stable motor loading for APP axonal transport, we found that APP velocity modes are relatively stable.

**Kinesin-1 reduction impairs retrograde transport**

To examine the effects of kinesin-1 reduction on the retrograde transport of APP, we first determined the retrograde behavior of APPYFP vesicles under control conditions. The observed mean retrograde segmental velocity was 0.87 μm/s (Figure 3A), similar to values reported for cytoplasmic dynein in vitro (King and Schroer, 2000; Reck-Peterson et al., 2006). However, retrograde segmental velocity distribution showed three modes, with velocities as high as 3.14 μm/s (Figure 3B and Table S1). As with anterograde transport, retrograde velocities showed high modal stability (Figure 3E). The observed retrograde run length was 7.08 μm (Figure S2J), and the calculated bulk retrograde run length was 13.2 μm. Longer-duration recordings confirmed the longer calculated retrograde run length (mean = 14.18 μm; SEM = 2.7 μm; n = 26 segments; 2 min of imaging; unpublished data).

**FIGURE 3:** Kinesin-1 reduction leads to impairment in retrograde APPYFP transport. (A) Distribution of retrograde segmental velocities. (B) Retrograde unweighted segmental velocity mode analysis for control shows three modes (cyan); see Table S1 for a definition of mode centers, spreads, and fractions of segment population. Superposition of modes is shown in red. A similar phenomenon is observed for khc<sup>8ex94</sup>/+ and klc<sup>8ex94</sup>/+ (see Table S1). Red line, superposition of modes. A similar phenomenon is observed for khc<sup>8ex94</sup>/+ and klc<sup>8ex94</sup>/+ (see Table S1). Red line, superposition of modes. A similar phenomenon is observed for khc<sup>8ex94</sup>/+ and klc<sup>8ex94</sup>/+ (see Table S1). Red line, superposition of modes. A similar phenomenon is observed for khc<sup>8ex94</sup>/+ and klc<sup>8ex94</sup>/+ (see Table S1). Red line, superposition of modes. A similar phenomenon is observed for khc<sup>8ex94</sup>/+ and klc<sup>8ex94</sup>/+ (see Table S1). Red line, superposition of modes.
Reduction in khc and klc gene dosage led to substantial impairment in retrograde transport. We found significant decreases in APPYFP segmental velocities (Figures 3C and S2I) and a major shift in occupancy (but not value) of velocity modes (Figures 3, D and E, and S5F and Table S1). In addition, there was a significant decrease in retrograde run lengths and significant increases in retrograde pause frequencies and durations (Figure S2, J–L). Thus kinesin-1 plays an important role in bidirectional transport, as proposed previously (Brady et al., 1990; Martin et al., 1999; Pilling et al., 2006; Kim et al., 2007; Colin et al., 2008). Although khc reductions produced more pronounced effects compared with klc, we observed that both subunits play a similar role, unlike previous work in which only KHC had a significant contribution (Ling et al., 2004).

We considered three explanations for the impairment in retrograde transport by kinesin-1 reduction: dynein vesicle loading, motor activation, and sampling bias.

If loss of kinesin-1 leads to reduced loading of cytoplasmic dynein on APP vesicles, as proposed previously (Ally et al., 2009), we should see evidence for this in our flotation assay. However, we noted that dynein heavy chain (DHC) protein levels in khc+/+ remained unchanged in the postnuclear supernatant (PNS) and exhibited only a minor (15%) decrease in the 8/35 fraction (Figure 2, D and E). Thus loss of kinesin-1 does not appear to significantly lower the amounts of cytoplasmic dynein on floated vesicles.

The hypothesis of dynein activation by kinesin (Ally et al., 2009) cannot be tested directly in our system. However, a prediction of this hypothesis is that kinesin overexpression will lead to increased dynein activation. Thus we overexpressed khc, klc, or both together (see Materials and Methods) and analyzed APPYFP movement. Western blot analysis confirmed the expected increase in KHC and KLC proteins in kinesin-1 overexpression genotypes (Figure 4, A–C). Movement analysis in these backgrounds showed a significant impairment in retrograde transport, with decreases in segmental velocities and run lengths, as well as increases in pause frequency and duration (Figures 4E and S2, I–L). These findings do not support the hypothesis that kinesin-1 activates cytoplasmic dynein. Intriguingly, we also observed a significant impairment in anterograde transport (Figures 4D and S2, A–D), even though there was a significant increase in the anterograde population fraction (Figure S2F).

Sampling bias attributes the observed impairment in retrograde transport to...
overall reduction in APP movement. The bias would result from fewer axonal retrograde vesicles within the observation window, resulting in an apparent inhibition of retrograde transport (Figure 4, F and G). Indeed, we observed a significant decrease in the total number of APPYFP vesicles in kinesin-1 reductions, an effect we observed for no other genotype (Figure 4H). Thus sampling bias alone may explain the apparent impairment of retrograde transport upon reduction of kinesin-1.

**Reduction of cytoplasmic dynein and dynactin subunits confirms modal stability is a generic feature of kinesin-1–driven APP transport**

Previous work in *Drosophila* embryos showed that impairment to DHC or the p150 subunit of dynactin disrupts anterograde movement (Gross et al., 2002). From this, dynactin emerged as a potential coordinating factor. Other work has proposed that dynactin promotes dynein attachment to vesicles and enhances retrograde motor processivity through p150 binding to microtubules (Karki and Holzbaur, 1999; King and Schroer, 2000; Schroer, 2004). However, more recent work demonstrated that cargo motility in vivo does not require p150 microtubule binding (Kim et al., 2007). In addition, loss of dynactin does not prevent dynein association with vesicles (Haghnia et al., 2007). Although disruption of the dynactin complex via arp1 deletion or p50 overexpression led to bidirectional impairment in axonal transport (Haghnia et al., 2007; Kwinter et al., 2009), the severity of the effects raised concerns about the specificity of the inhibition. Thus we further probed models of motor protein coordination and attachment by evaluating the effects of reduction in cytoplasmic dynein and dynactin on APP transport (see Materials and Methods). For cytoplasmic dynein, we studied two null point mutation alleles for DHC, dhc64c<sup>1412</sup> and dhc64c<sup>1413</sup> (see Materials and Methods), two null point mutations for dynein IC (DIC), dic<sup>1</sup> and dic<sup>2</sup> (Boylan and Hays, 2002), and one deletion mutation of the dynein light chain (DLC), rob<sup>h</sup> (Bowman et al., 1999). Western blot analysis in dynein and dynactin heterozygotes showed the expected ~50% reduction in protein levels (DHC: Figure 5, A and B; DIC: Figure S3A; DLC: Figure S3B; dynactin: Figure S4, A and B).

For dynein, dhc reduction resulted in an expected overall impairment of many features of retrograde transport but enhancement of a number of anterograde parameters (Figures 5, C–E, S3, C–J, and SSC, and Table S1), including significant increases in anterograde run length (Figure S3E) and average segmental velocity (Figure S3C), as well as shifts in the relative contribution of higher modes to the anterograde velocity distribution (Figures S3E and SSC and Table S1). This suggests that DHC acts as an inhibitor of anterograde transport. DLC behaved in similar manner, although the overall effect on anterograde was milder (e.g., significant increase in anterograde run length [Figure S3E], but not segmental velocity [Figure S3C]). In contrast, dic reduction not only led to impairment in retrograde transport but also to strong impairment in anterograde transport (Figures 5, C and D, and S3, C–J). As with kinesin, velocity modal stability was maintained in dynein reductions (Figure 5F). Thus DHC/DLC appear to act as inhibitors of anterograde movement, while DIC acts as a promoter.

In general, dynactin reduction led to bidirectional impairment of APP transport (Figures 5, G and H, and S4, C–F, and Table S1), although some variability was observed among the subunits. For dmn (p50 dynamitin), there was a mild decrease in mean retrograde segmental velocity but no significant change in anterograde velocity (Figure 5, G and H). However, there was a significant increase in both pause frequency and duration (Table S1) and in the fractions of the population of vesicles with anterograde movement (Figure S4, G–J). Perturbation of G{p150<sub>Glu26</sub>} and grid (Arp1) caused stronger transport impairment in terms of reduced segmental velocities and run lengths (Figures 5, G and H, and S4, C–F). However, the impairment was less pronounced in terms of increased pause frequencies and durations (Table S1). As with kinesin and dynein, velocity modes were maintained in dynactin reductions (Figure 5I). Overall, these data confirm that dynactin promotes anterograde and retrograde transport, although the transport impairment associated with perturbation in these dynactin subunits is milder than is observed for kinesin-1 and cytoplasmic dynein. In addition, these data support the notion that modal stability is a generic feature of APP transport.

**Stimulation of anterograde transport by overexpression of DIC**

Cytoplasmic dynein has been proposed as an activator of kinesin-1 (Martin et al., 1999; Ling et al., 2004; Pilling et al., 2006; Kim et al., 2007). Previous biochemical data established that the only detectable interactions between kinesin-1, cytoplasmic dynein, and dynactin protein complexes are mediated by DIC. Interactions were reported between DIC and p150<sub>Glud</sub> (Vaughan and Vallee, 1995) and DIC-KLC (Ligon et al., 2004) and appear to involve similar regions of DIC (Vaughan and Vallee, 1995; Ligon et al., 2004; Schroer, 2004). Previous work, however, did not distinguish whether KLC, dynactin, or both mediate the interaction between endogenous kinesin and dynein, since the in vitro work demonstrating an interaction between KLC and DIC used recombinant protein for the affinity chromatography assay (Ligon et al., 2004). Thus we performed RNA interference (RNAi) experiments in S2 cells to test whether KLC binds to DIC in vivo (see Materials and Methods). We observed that KLC mediates binding of the KHCl/KLC kinesin-1 tetramer to DIC (Figure 6A). Hence, it is possible that motor protein activity on APP vesicles is controlled by a competition between p150<sub>Glu26</sub> and KLC for binding to DIC. Consistent with this idea, we noted that among the dynein reduction subunits, DIC reduction caused by far the strongest and most consistent impairment to bidirectional transport.

If DIC controls the balance between anterograde and retrograde transport by activating kinesin-1, overexpression of DIC would stimulate anterograde transport. To test this, we overexpressed DIC protein (see Materials and Methods; Figure S3K). This led to significant increases in both anterograde vesicle fraction and run length (Figure S3, E and H), as well as significant decreases in the fraction of stationary and reversing vesicles (Figure S3, G and J). In addition, the mean anterograde velocity increased significantly (Figure S3C), due to a systematic shift of each of the three velocity distribution modes toward higher center values (Figures 6D and SSD and Table S1). It should be noted that this was the only genetic condition for which we observed significant shifts in the mode centers. Taken together, these findings indicate that DIC acts as a promoter of kinesin-1 activity.

**DISCUSSION**

In this study, we used modest genetic alterations of kinesin-1, cytoplasmic dynein, or dynactin to probe mechanisms of axonal transport in vivo. Computer-assisted quantification of large numbers of APPYFP vesicles allowed us to achieve sufficient sensitivity for robust interpretation of the mild phenotypes induced by manipulations that maintain full organismal viability and are devoid of major uncontrolled side effects. We discovered several novel features of APP transport. First, vesicle movement driven by kinesin-1 in vivo has crucial differences from in vitro: while in vitro cargo velocity is independent of the number of motors, the velocity of kinesin-1-driven...
FIGURE 5: Reduction of cytoplasmic dynein and dynactin subunits confirms that modal stability is a generic feature of kinesin-1–driven APP transport. (A) Generation of dhc null alleles for evaluation of APP transport. Protein extracts were either treated with UV-vanadate (+) or left untreated (−) and assayed by Western blot (see Materials and Methods). The anti-DHC antibody recognizes endogenous DHC (the DHC-3HA transgene product) as well as the LUV fragments resulting from UV cleavage of the endogenous and transgenic DHC. Tagged and untagged LUV fragments are resolved as a doublet. Only a single LUV band (corresponding to the larger, tagged fragment) is recognized in the rescued Df/g genotype, consistent with the absence of endogenous DHC protein expression. Similar results are seen with the DHC allele 4-19 (far right). (B) Western blot analysis of cytoplasmic dynein reduction genotypes probed for DHC shows an ∼50% reduction in dhc64c4-19/+ and ∼70% in dhc*p4163/+ . Significant reduction in DHC is also observed in dlc reduction backgrounds. (C) Reduction in dhc and dlc shows no significant effect on anterograde duration-weighted segmental velocities (mean μm/s ± SD): control, 1.09 ± 0.58; dhc64c4-19/+ , 1.14 ± 0.66; dhc*p4163/+ , 1.06 ± 0.52; robl/+ , 1.07 ± 0.62. Reduction in dlc shows significant impairment in the anterograde duration-weighted segmental velocities (mean μm/s ± SD): control, 1.09 ± 0.58; dlc/-/+ , 0.91 ± 0.60; dlc/-/+ , 0.92 ± 0.55. (D) Reduction in dhc, dlc, and dlc alleles leads to a significant decrease in retrograde duration-weighted segmental velocities (mean μm/s ± SD): control, 1.02 ± 0.41; dhc64c4-19/+ , 0.93 ± 0.50; dhc*p4163/+ , 0.89 ± 0.38; robl/+ , 0.93 ± 0.53; dlc/-/+ , 0.78 ± 0.48; dlc/-/+ , 0.68 ± 0.37. (E) The anterograde segmental velocity distribution of dhc64c4-19/+ has three modes (cyan) with a shift toward occupancy of modes 2 and 3 (see Table S1; the control distribution is shown in yellow for direct comparison). Similar behavior is found for dhc*p4163/+ . (F) Linear regression of retrograde mode centers assembled for dynein reduction genotypes shows centers that follow a ratio of ∼1:2:4. (G) Reduction in GlI (p150) and grid (Arp1) subunits of dynactin leads to a significant decrease in mean anterograde duration-weighted segmental velocity (mean μm/s ± SD): control, 1.02 ± 0.41; dmn5499/+ , 0.93 ± 0.44; GlI/+ , 0.88 ± 0.42; grid/+ , 0.85 ± 0.44.
FIGURE 6: DIC stimulates anterograde APPYFP transport. (A) RNAi pulldown experiments in S2 cells demonstrate that KLC mediates binding of the KHC/KLC kinesin-1 tetramer to DIC in vivo. (B) DIC overexpression has no significant effect on mean anterograde duration-weighted segmental velocity compared with control (mean μm/s ± SD): control, 1.09 ± 0.58; GEN-DIC, 1.14 ± 0.67. However, GEN-DIC leads to a significant increase in mean anterograde unweighted segmental velocity and several other parameters of anterograde transport (Figure S3, C–J). (C) DIC overexpression has no significant effect on mean retrograde duration-weighted segmental velocity compared with control (mean μm/s ± SD): control, 1.02 ± 0.41; GEN-DIC, 1.03 ± 0.52. (D) The anterograde segmental velocity distribution of GEN-DIC has three modes (cyan) with a significant shift of mode centers toward higher values (see Table S1 and Figure S5D for a direct comparison with modes of the control distribution). Superposition of modes is shown in red. (E) The retrograde segmental velocity distribution of GEN-DIC has three modes (cyan; see Table S1 and Figure S5H for a direct comparison with modes of the control distribution). Superposition of modes is shown in red. (F) Schematic of a proposed model for APP transport under control, kinesin-1 overexpression, and DIC overexpression backgrounds (A = anterograde; R = retrograde; S = stationary). Excess kinesin-1 sequesters DIC from association with vesicle-bound KLC. This results in "decreased activation" of anterograde transport. This sequestration of DIC may also result in inhibition of retrograde transport via reduced vesicle-bound dynein or dynactin. Under excess DIC conditions, anterograde movement can be stimulated by either increased vesicle-bound DIC (i.e., cytoplasmic dynein complex) or soluble DIC associating with vesicle-bound KLC.
APP vesicles in vivo depend strongly on the number of motors. This deviates fundamentally from observations reported for lipid droplet movement (Shubeita et al., 2008). Second, APP vesicles exhibit relatively stable modal velocities, suggesting the presence of stable motor assemblages on vesicles. Third, our data suggest a new and testable motor competition–coordination model of axonal transport in which direction and velocity of cargo movement are controlled at least in part by DIC.

**Does the amount of kinesin-1 on APP vesicles control movement parameters?**

Considerable in vitro biochemical and biophysical data indicate that kinesin-1 is highly processive. It has been proposed that the processivity originates in the strong binding of the motor head for a large fraction of the total mechanochemical cycle and the tight coordination of the binding and release of the two heads (Spudich, 1990; Howard, 2001; Gennerich and Vale, 2009). The high processivity of kinesin-1 has been postulated to be essential for long-distance vesicle movement in vivo, as it reduces the probability of cargo dissociation from microtubules.

A key prediction for processive motor transport is that the run length of cargo correlates with the number of motors bound, while the velocity is independent. Although recent work suggests that velocity of droplet movement in Drosophila embryos is indeed uncorrelated with the amount of active kinesin-1 (Shubeita et al., 2008), multiple studies have reported kinesin-1 velocities that are too fast to easily reconcile with the in vitro rate (Howard, 2001; Kural et al., 2005). This raised the possibility that both velocity and run length of kinesin-1 cargoes in vivo are dependent on the amount of available motor, a prediction fully supported by our analyses of the velocities and run lengths of kinesin-1-driven APP vesicles.

There are at least three possible explanations for this finding. First, the motor could be regulated to have a faster ATPase rate in vivo. However, this explanation does not readily account for the observation that run length and velocity of APP vesicles are correlated in vivo. Second, kinesin-1 on APP vesicles might be regulated or modified in vivo, such that the rate-limiting, strongly bound state time during the mechanochemical cycle is reduced. As a consequence, vesicle movement would show strong correlation of motor number with both run length and velocity, and run length or pause frequency would strongly depend on velocity (Spudich, 1990). Although a molecular explanation for this type of kinesin-1 behavior is lacking, recent work suggests that regions adjacent to the motor domain may control motor coordination (Crevenna et al., too). Notably, previous work on the biophysical behavior of the coil 1 domain of KHC suggests that this structured alpha-helical coiled-coil has the unusual ability to melt at physiological temperature (de Cuevas et al., 1992). Melting of this coiled-coil in vivo could reduce coordination between the heads within the kinesin-1 dimer, diminishing processivity and increasing velocity and run lengths generated by multiple kinesin-1 motors. A third possibility is that the strongly bound state time of kinesin-1 might not change in vivo, but the coiled-coil stalk domain might stretch like a spring, such that the velocity of a single kinesin-1 motor is no longer rate limiting for assembly of multiple motors on a vesicle. This possibility would also account for the observed correlation of kinesin-1 amount and run length of APP vesicles. Further biophysical work will be necessary to distinguish among these possibilities.

Our results also suggest that the assemblage of motors and their activation states on APP vesicles stay relatively constant for the majority of vesicles, even over the duration of a pause. Such “memory” on the timescale of seconds was also noted for lipid droplets (Welte et al., 1998).

**Movement of APP vesicles follows a multimodal velocity distribution**

Although multimodal velocity distributions of active cargo transport have been identified previously in vitro and in vivo (Hill et al., 2004; Levi et al., 2006; Gagliano et al., 2010), we made several new findings in this study. First, we analyzed and inferred multimodal velocity distributions using different computational and statistical analysis methods and assumptions. Previous studies (Hill et al., 2004; Levi et al., 2006; Gagliano et al., 2010) assumed a model in which multiple motors share their mechanical load. Consequently, it was assumed that higher-velocity modes are multiples of the lowest mode, presumably generated by a single motor, and that each velocity mode has the same spread (i.e., variance). We made no such assumptions in our analysis. Instead, we reason that multiple factors may be involved in determining velocities of actively transported cargoes and thus prefer an approach that permits varying spread within each mode and varying separation between neighboring modes. Our analysis consistently identified three velocity modes that follow a ratio of ∼1:2:4 in both anterograde and retrograde directions. The identified velocity modes we identified are not equally spaced and have different variances, with higher velocity modes associated with larger variances. Second, we identified how different genetic manipulations of individual units of the molecular motor machinery may differentially shift relative contributions of individual modes. Our analysis consistently returned three velocity modes that follow a ratio of 1:2:4 in all genetic backgrounds we analyzed. This provides further support for the analysis method we used. Third, we found that velocities of individual APP vesicles are relatively stable based on computational analysis of transition probabilities between different modes. Such analysis provides new insights into the dynamic behavior of individual cargoes.

**The in vivo behavior of APP vesicles is inconsistent with most current models for control of vesicle movement**

In agreement with previous work on lipid droplets, melanosomes, and other nonneuronal cargoes, our data do not support simple tug-of-war or exclusionary presence models and are most consistent with coordination models (Gross, 2004; Kural et al., 2005). For example, an exclusionary presence model for APP axonal vesicles predicts that reduction of kinesin-1 motors should have either no effect or increased retrograde transport, while reduction of dynein motors should have no effect or increased anterograde transport. Similarly, increased kinesin-1 is predicted to have either no effect or increased anterograde transport and decreased retrograde transport. For simple tug-of-war models, the prediction is that simultaneous attachment and activity of opposing motors would generate net APP movement in one direction or another, depending on the relative number of each type of motor attached to a vesicle (Mueller et al., 2008; Soppina et al., 2009). Thus reduction of either kinesin-1 or cytoplasmic dynein would be predicted to uniformly stimulate movement in the opposite direction, which is inconsistent with many of our observations. A more sophisticated tug-of-war model (Ally et al., 2009) proposed that dynein activity is needed to activate kinesin-1 and vice versa. Indeed, in agreement with previous reports, kinesin reductions seemed to inhibit dynein-driven movements (Brady et al., 1990; Martin et al., 1999; Pilling et al., 2006; Kim et al., 2007; Colin et al., 2008). However, although we cannot completely exclude direct effects of kinesin-1 reduction on cytoplasmic dynein activity, sampling bias is a compelling explanation for the impaired
dynein-driven retrograde transport upon kinesin-1 reduction (see Figure 4, F–H). More importantly, our finding that reductions in DHC enhance anterograde movement is inconsistent with the model proposed by Ally and coworkers (Ally et al., 2009).

A computational model of motor-driven transport was recently proposed (Muller et al., 2008) in which multiple kinesin and dynein motors interact indirectly via their nonlinear, force-dependent unbinding rates. This yields a nonintuitive coordination mechanism whereby vesicle movement in a specific direction is driven predominantly by one type of motor. We tested several predictions of this model using our data. First, when the plus end–directed and minus end–directed motors are asymmetric, the model predicts that anterograde and retrograde vesicle movement each follow a single peak–velocity distribution. However, our APPYFP data reveal that distributions of both anterograde and retrograde velocities are much wider and exhibit multiple modes. Second, the model proposes that weak motors with a small ratio of stall-to-detachment forces drives stationary cargoes classified under the category of “no motion.” Consequently, the model predicts that a significant increase in the fraction of stationary cargoes results from increases in both weak plus end–directed and minus end–directed motors. In fact, significant increases in the fraction of stationary cargoes are observed in many of our reduction genotypes. For example, KHC reductions (khc8/ and khc20/+) cause a twofold increase in the fraction of stationary cargoes (Figure S2E). This would require a concurrent increase in the fraction of “weak” dynein motors, which seems at odds with the basic model assumption that changes to kinesin do not alter the inherent properties of dynein motors and vice versa. Finally, although changing the parameters of the motors can account for changes to stall force, run length, pause time, and velocity caused by different dynein mutations in the lipid droplet system, this model cannot account for the changes we observe for APP vesicles in the population distribution of different types of cargoes (anterograde, retrograde, stationary, and reversing). For example, the model cannot explain why there is a substantial fraction (∼40%) of stationary APPYFP vesicles in the control group and why there are different fractions of anterograde (∼32%) and retrograde (∼18%) cargoes. These arguments suggest that mechanical coordination alone cannot fully account for the behavior of APP axonal transport.

A competition–coordination mechanism regulates axonal transport of APP vesicles

In a typical coordination model, vesicles bind to both kinesin-1 and cytoplasmic dynein simultaneously, and a coordination process on the vesicle surface ensures that both types of motors are not active at the same time (Gross, 2004). The predictions of this model are ambiguous, given the lack of a defined coordination mechanism. However, if kinesin and dynein themselves are part of the coordination mechanism, this model would predict that reductions of either kinesin-1 or cytoplasmic dynein might impair movement in both directions. For vesicle transport in the lipid droplet system, it was argued that a coordination model best accounts for the experimental observations. Our data constrain the type of coordination model that can be proposed. We find that: 1) reducing kinesin-1 does not decrease cytoplasmic dynein levels on floated vesicles, and hence kinesin-1 is not required for cytoplasmic dynein association with vesicles; 2) overexpression of kinesin-1 does not stimulate dynein-driven retrograde transport, suggesting that kinesin-1 binding to DIC does not stimulate dynein activity; 3) sampling effects in kinesin-1 reduction may account for decreases in retrograde movement; and 4) modest overexpression of DIC leads to a significant increase in retrograde run length but a much greater stimulatory effect is observed for anterograde movement in terms of distribution and occupancy of velocity modes toward faster-moving modes, longer anterograde run lengths, and a large increase in the ratio of anterograde to retrograde movement (a shift from 1.8:1 to 2.3:1). Thus a formal genetic interpretation of our results implicates DIC as an activator of kinesin-1–mediated anterograde transport. Combining these genetic findings with biochemical evidence for the ability of DIC to bind p150Glued and KLC in similar regions (Ligon et al., 2004) leads to a simple model in which the activity of kinesin-1 and cytoplasmic dynein is coordinated by competition of dynactin and kinesin-1 for binding to DIC, thereby controlling the direction of movement (Figure 6F). This model proposes that the binding of dynactin to DIC promotes retrograde transport and binding of DIC to KLC promotes anterograde transport. Excess soluble kinesin-1 is predicted to compete with vesicle-bound dynactin for DIC binding, resulting in impaired activation of cytoplasmic dynein. Similarly, excess kinesin-1 is predicted to compete with vesicle-bound KLC for DIC binding, causing impaired kinesin-1 movement. Indeed, our data exhibit reduced bidirectional transport in kinesin-1 overexpression (Figures 4, D and E, and S2, A–D and I–L). On the other hand, the model predicts that excess soluble DIC binds preferentially to KLC on APP vesicles, resulting in enhanced anterograde transport bias. Indeed, for vesicles in the GEN (genomic)-DIC genotype, we found a marked increase in anterograde population percentage, faster average segmental velocities, increase in occupancy and mean value of mode 3 (fastest mode), and extended anterograde run lengths (Figure S3, G, C, D, and E, respectively, and Table S1). Thus DIC activates anterograde transport and biases axonal transport in the anterograde direction. The precise mechanism for this bias in GEN-DIC is an interesting question warranting further investigation. Intriguingly, we also observed that overexpression of DIC increased bidirectional pause frequency and duration (Table S1). Because this occurred in both anterograde and retrograde directions and without reduction in duration-weighted segmental velocities (Figure 6, B and C), it is possible that increased density of non-APP vesicles (obstacles) along the axon indirectly interferes and/or temporally stalls vesicle movement in this genotype. Taken together, our data provide strong quantitative support for a competition–coordination mechanism for the in vivo axonal transport of APP in which DIC–KLC interactions define a switch in the control of retrograde and anterograde movement. Previous overexpression studies in nonneuronal cells were interpreted to mean that excess DIC poisons retrograde transport (King et al., 2003). However, unlike our live-cell measurements, previous fixed-cell analyses could not distinguish inhibition of retrograde transport from stimulation of anterograde transport or a combination of the two. Still, we cannot completely exclude that some aspects of our data might represent indirect effects of protein overexpression on transport (i.e., due to aberrant transport/sequestration of known or unknown regulators of motor proteins). For example, kinesin-1 overexpression did not yield faster anterograde velocities, indicating that additional regulatory factors are required (Gindhart et al., 2003; Blasius et al., 2007). Further investigations will be needed to test the many detailed predictions of our model. In addition, a search for various proteins interacting with p150Glued, DIC, and KLC may augment our understanding of how the interactions between DIC, dynactin, and/or KLC may be modulated.

Finally, the rich biological diversity observed within and across microtubule motor families, coupled to the unique protein constitution of different cargo types, as well as multiple regulators of axonal transport, raises the possibility that no single universal mechanism may emerge for axonal transport. Indeed, controversies in the literature...
may reflect this heterogeneity. These differences highlight the importance of employing well-characterized vesicles in the study of axonal transport in vivo. Our ability to rescue axonal transport defects seen in a variety of neurodegenerative diseases affecting humans will likely reflect this heterogeneity due to instability (Saxton et al., 1996; Hurd and Saxton, 1996; Hurd et al., 1996; Brendza et al., 2000). For kinesin-1 reduction studies, three alleles were used: khc4163a/CyO, khc4163a/CyO, and klc8ex94/TM6B. The khc alleles are null point mutants whose translated fragments are thought to be degradable due to instability (Saxton et al., 1991; Hurd and Saxton, 1996; Hurd et al., 1996; Brendza et al., 2000). For dhhc4164a/+;TM6B, dhhc4164a/+;TM6B, dcl/;FM7GFP, dcl/;FM7GFP, and rob/;CyO. The dhhc4164a/+ allele has an internal 122–base pair deletion of nucleotides 5750–5871 (with respect to the Dhc64C cDNA), which is predicted to remove P-loop 1 and lead to premature termination of translation upstream of P-loops 2–4. Western blot analysis of these mutations confirms that dhhc4164a/+ and dhhc4164a/+ are null or near-null alleles (Figure S5A). The dcl alleles have been characterized as loss of function, with dcl being functionally stronger than dcl (Boylan and Hays, 2002). The dcl allele rob/+ is a deletion allele spanning the entire robl gene (Bowman et al., 1999). For crosses involving dhc or dlc, we followed the procedure used for kinesin-1. For dcl crosses, UAS-APPYFP/Y; SG26.1Gal4/B3 animals were crossed to dcl/;FM7GFP or dcl/;FM7GFP, and a fluorescence microscope was used to select female L3 larvae negative for GFP—these animals are UAS-APPYFP/dcl; SG26.1Gal4/+.

For dynactin reduction studies, three alleles were used: grid/TM6B, dmn5497/CyO, and Gl'/In(3L)U, d' (BL-2394). The grid allele has been characterized as a loss-of-function point mutant (Haghnia et al., 2007). The dmn5497 allele, kindly provided by Jason Duncan (Williamette University, Salem, OR), is also thought to be a loss-of-function point mutant (unpublished observations). The Gl' allele is thought to be dominant negative. As with kinesin-1, UAS-APPYFP, pin89B/V3 virgins were crossed to grid/TM6B, dmn5497/CyO, or Gl'/In(3L)U, d' (BL-2394 Bloomington stock). Selected UAS-APPYFP/Y; dynactin/V3 males were then crossed to SG26.1Gal4 females, and non-tubby L3 females were selected for imaging.

For kinesin-1 overexpression experiments, we used GEN-KLC/CyO; TM3/TM6B and sp/CyO; GEN-KHC stocks (Saxton et al., 1991; Gindhart et al., 1998). To achieve overexpression of both kinesin-1 subunits, we crossed GEN-KLC; GEN-KHC females to UAS-APPYFP/Y; SG26.1Gal4/B3 males. UAS-APPYFP/X; GEN-KLC/+; GEN-KHC/SG26.1 animals were used for imaging.

For cytoplasmic dynein overexpression experiments, we used a GEN-DIC stock (Boylan and Hays, 2002). In brief, UAS-APPYFP/Y; GEN-DIC/B3 males were generated and crossed to SG26.1 virgins. UAS-APPYFP/X; GEN-DIC/+; SG26.1/+ L3 larvae were then selected for imaging.

**Microscopy**

L3 larvae were dissected and imaged on a 3 cm² Sylgard platform using Ca²⁺-free medium consisting of NaCl (128 mM), ethylene glycol tetraacetic acid (EGTA; 1 mM), MgCl2 (4 mM), KCl (2 mM), HEPES (5 mM), and sucrose (36 mM). Dissected animals were inverted onto a coverslip and imaged using a Nikon Eclipse TE2000-U inverted microscope with a CoolSNAP HQ camera (Roper Scientific, Surrey, BC, Canada) and a 100×/1.40 numerical aperture (NA) oil objective. Images were collected at 10 frames/s with 100 ms exposure for 15 s under the control of MetaMorph software (MDS Analytical Technologies, Sunnyvale, CA). For each genotype, four time-lapse movies were collected for each animal (10 animals were imaged for analysis).

To characterize Dhc64C gene mutations, we crossed balanced females carrying the γ4163a or 4-19 mutations to males expressing the Dhc-3HA transgene and a deficiency (Df) that removes the Dhc64C gene. Progeny of genotype Df/y or Df/4-19 survive only in the presence of the transgene. Protein extracts from all the female progeny classes were either treated with UV-vanadate (+) or left untreated (−) and then assayed by Western blot. The blot was probed with an anti-DHC antibody that recognizes endogenous DHC (the DHC-3HA [hemagglutinin] transgene product) and the LUV (light ultraviolet cleavage fragment) fragments resulting from UV cleavage of the endogenous and transgenic DHC. Tagged and untagged LUV fragments are resolved as a doublet. Only a single LUV band is recognized in rescued Df/y or Df/4-19 flies, corresponding to the higher-molecular-weight tagged fragment. This is consistent with the absence of endogenous DHC protein expression.

**Protein analysis of kinesin-1, cytoplasmic dynein, and dynactin genetic reduction and overexpression backgrounds was performed using Western blot and quantified with an Odyssey system (Li-Cor Biosciences, Lincoln, NE). In brief, 10 animals were homogenized in 100 μl 2X LDS sample buffer (Invitrogen, Carlsbad, CA) with 2% β-mercaptoethanol. Samples were heated for 10 min at 98°C and centrifuged for 5 min at 14,000 rpm. The supernatant was then removed, and protein concentration was measured using the Bio-Rad DC protein assay (Hercules, CA). Samples were loaded on pre-cast 4–12% Bis-Tris gels (Invitrogen), and See-Plus 2 (Invitrogen) was used as the molecular-weight marker. Transfer to nitrocellulose membranes was done at 4°C using a wet system and buffer consisting of 25 mM Tris base, 190 mM glycine, and 20% methanol. Membranes were incubated in primary antibody overnight at 4°C. Incubation in secondary antibody (1:5000; Odyssey) was 2 h at room temperature. Quantification was done using Odyssey software (Li-Cor, Lincoln, NE).

**Antibodies**

Anti-KHC (Cytoskeleton, Denver, CO), anti-KLC (Goldstein lab), anti-DHC (PIH4; McGrl and Hays, 1997), anti-DIC (Chemicon International, Temecula, CA), anti-ROBL (Bowman et al., 1999), anti-Arp1

**MATERIALS AND METHODS**

**Genetics**

*Drosophila* stocks were maintained at room temperature or at 25°C, except when crosses were set up at 29°C for imaging of L3 larvae. UAS-APPYFP (X) controls carry the human APP695 transgene tagged with YFP and inserted in the X chromosome. Only female L3 larvae were used for imaging. In brief, L3 larvae expressing UAS-APPYFP arose from crosses between UAS-APPYFP males and SG26.1Gal4 females. Male animals did not express UAS-APPYFP and thus provided an internal control. Motor gene reductions were obtained by first generating a UAS-APPYFP; pin89B/B3 stable stock (the B3 chromosome carries T(2;3) CyO, TM6B with the dominant markers Hu, Tb, and Cyo).

For kinesin-1 reduction studies, three alleles were used: khc4163a/CyO, khc4163a/CyO, and klc8ex94/TM6B. The khc alleles are null point mutants whose translated fragments are thought to be degraded due to instability (Saxton et al., 1991; Hurd and Saxton, 1996; Hurd et al., 1996; Brendza et al., 2000). The klc allele is a deletion allele spanning the entire klc gene in chromosome 3 (Gindhart et al., 1998). UAS-APPYFP; pin89B/V3 females were crossed to khc/CyO, khc/CyO, or klc8ex94/TM6B males. Selected UAS-APPYFP/Y; kinesin/B3 males were then crossed to SG26.1Gal4 females. Non-tubby L3 females were selected for imaging.

For cytoplasmic dynein reduction studies, five alleles were used: dhhc4164a/+;TM6B, dhhc4164a/+;TM6B, dhhc4164a/+;TM6B, dhhc4164a/+;TM6B, and dhhc4164a/+;TM6B, dcl/;FM7GFP, dcl/;FM7GFP, and rob/;CyO. The dhhc4164a/+ allele has an internal 122–base pair deletion of nucleotides 5750–5871 (with respect to the Dhc64C cDNA), which is predicted to remove P-loop 1 and lead to premature termination of translation upstream of P-loops 2–4. Western blot analysis of these mutations confirms that dhhc4164a/+ and dhhc4164a/+ are null or near-null alleles (Figure S5A). The dcl alleles have been characterized as loss of function, with dcl being functionally stronger than dcl (Boylan and Hays, 2002). The dcl allele rob/+ is a deletion allele spanning the entire robl gene (Bowman et al., 1999). For crosses involving dhc or dlc, we followed the procedure used for kinesin-1. For dcl crosses, UAS-APPYFP/Y; SG26.1Gal4/B3 animals were crossed to dcl/;FM7GFP or dcl/;FM7GFP, and a fluorescence microscope was used to select female L3 larvae negative for GFP—these animals are UAS-APPYFP/dcl; SG26.1Gal4/+.

For dynactin reduction studies, three alleles were used: grid/TM6B, dmn5497/CyO, and Gl'/In(3L)U, d' (BL-2394). The grid allele has been characterized as a loss-of-function point mutant (Haghnia et al., 2007). The dmn5497 allele, kindly provided by Jason Duncan (Williamette University, Salem, OR), is also thought to be a loss-of-function point mutant (unpublished observations). The Gl' allele is thought to be dominant negative. As with kinesin-1, UAS-APPYFP, pin89B/V3 virgins were crossed to grid/TM6B, dmn5497/CyO, or Gl'/In(3L)U, d' (BL-2394 Bloomington stock). Selected UAS-APPYFP/Y; dynactin/V3 males were then crossed to SG26.1Gal4 females, and non-tubby L3 females were selected for imaging.
(Haghnia et al., 2007), anti-p50 (Duncan and Warrior, 2002), and anti-synaptin (8C3; Developmental Studies Hybridoma Bank, University of Iowa, Iowa City, IA) were used as described or as recommended by the manufacturer. Many commercially available anti-p150 antibodies were tested, but none recognized Drosophila p150. Additionally, RNAi pulldown experiments utilized polyclonal anti-glued antibody (Waterman-Storer and Holzbaur, 1996) and a polyclonal pan-anti-KLC antibody (Khodjakov et al., 1998).

Membrane flotation assay
After raising yw and khc1+ animals at 25°C, we collected ~7.5 ml of adult flies in a 50-ml conical tube and immediately snap-froze them in liquid nitrogen. The tubes were shaken to separate heads from bodies, and fly heads were collected using a sieving system (Slemmon et al., 1982). Heads were then homogenized in 0.75 ml homogenization buffer composed of 8% sucrose plus 1 mM imidazole. Debris was removed by centrifugation at 1500 x g for 10 min at 4°C, and the PNS was collected. The pellet was then resuspended in 0.75 ml homogenization buffer and centrifuged for 5 min at 1500 x g. The supernatant was collected and combined with the PNS, which was centrifuged at 10,000 x g for 5 min to remove mitochondria. The resulting supernatant was carefully removed so as not to disturb the pellet and was mixed with 62% sucrose to a final concentration of 40% sucrose (“input”). The input was bottom-loaded in an 11 x 34 mm Beckman polycarbonate tube and overlaid with two 0.7 ml cushions of 35 and 8% sucrose. Tubes were spun at 200,000 x g for 2 h at 4°C. The 8/35% interface, which contains light membrane-bound organelles and associated proteins (Haghnia et al., 2007) was collected and assayed by SDS–PAGE and Western blotting.

KLC-RNAi pulldown assay
KLC RNAi expression in S2 cells was achieved by a double-stranded RNA preparation with a Megascript T7 kit (Applied Biosystems, Bedford, MA) and the following primers: sense 5’-t7-CTCGAAGCCTTTGCCAGTG-3’ and antisense 5’-t7-AGAGTACGGGACCAGCGCTG-3’. An N-terminal glutathione S-transferase (GST)-fusion protein of full-length Drosophila DIC (“GST-74”) was expressed in Escherichia coli and bound to glutathione Sepharose 4B beads (GE Healthcare, Waukesha, WI). Control S2 cells and RNAi-treated cells were homogenized in PEMU buffer (100 mM PIPES, 5 mM MgOAc, 5 mM EGTA, 0.5 mM EDTA, 0.9 M glycerol, 1 mM dithiothreitol; pH 6.9) with protease inhibitors. TX-100 0.1% was added, and the cells were centrifuged at 12,000 x g for 10 min at 4°C. Supernatants were preclarified against washed beads and subsequently incubated with GST-74 beads (or control beads bound to GST alone) for 2 h at 4°C. Beads were washed three times, and bound proteins were eluted into SDS–PAGE sample buffer for analysis by Western blot using 6–10% gradient gel and 15 μg protein.

Single-particle tracking and computational analysis of APPFYF vesicle movement
APPFYF vesicles in time-lapse movies were detected as particles (Ponti et al., 2003; see Supplemental Movie S2). Their full trajectories were recovered using customized software based on a modification of the single-particle tracking technique described previously (Yang et al., 2005). For vesicles crossing one another during bidirectional transport, trajectories were first resolved using a global bipartite linear assignment algorithm that minimizes total variations of intensity, velocity, and direction of movement for each vesicle. The output of this step contains mostly trajectory segments, so an additional process was performed to link these segments into full trajectories using a multiple hypothesis-testing algorithm (Blackman, 1999). A manual process was subsequently used to recover trajectories the software was unable to recover. Finally, all recovered trajectories were individually inspected, and errors were corrected. This process is able to recover more than 95% of all vesicle trajectories that fully reside in the microscope field of view.

For each genotype, individual cargoes were classified as being stationary, anterograde, retrograde, or reversing. Cargo trajectories of each genotype were then analyzed by calculating overall distribution of cargo population, as well as individual cargo velocity, run length, pause, and reversal (see Supplemental Material). Data analysis was conducted using customized software written in MATLAB (MathWorks, Natick, MA) and C++. Segmental velocities were defined as the mean velocity of a trajectory segment uninterrupted by a pause, reversal, or movie termination event. Duration-weighted segmental velocity evaluates the average velocity behavior that vesicles exhibit per time spent moving. Unweighted segmental velocity reports the average velocity without considering the duration of different trajectory segments. Estimated run length was calculated by dividing the mean segmental velocity by the total pause frequency.

Statistical analysis
Data were first checked for normality using three different tests in the most extreme package of R: the Lilliefors test, the Anderson-Darling test, and the Shapiro-Francis test. For those that followed normal distributions, their means and distributions were compared using pairwise two-sample, two-sided Student’s t tests and Kolmogorov-Smirnov tests, respectively. For nonnormal distributions, nonparametric tests, including permutation tests (Moore and McCabe, 2005) and Wilcoxon tests, were used (see Supplemental Material).

Modes in cargo velocities were identified by model-based clustering of the collection of all cargo segmental velocities using the mclust package of R (Fraley and Raftery, 2002). For each individual cargo, the mode of its segmental velocities can be determined using standard pattern classification techniques (Theodoridis and Koutroubas, 2009). Specifically, we compared the segmental velocity modes of each individual cargo in three cases: 1) before and after a pause; 2) before and after a reversal; and 3) on initiation and termination of a track segment. By repeating this analysis for all individual cargoes, we calculated the ensemble probabilities of transition between modes (i.e., probabilities of cargoes remaining in the same segmental velocity mode and switching between different modes). To examine whether individual cargoes stably maintain their velocity modes, we calculated the theoretical mode transition probabilities if cargoes were to randomly switch between different velocity modes (see Supplemental Material).

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