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Clathrin packets move in slow axonal transport and deliver functional payloads to synapses

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SUMMARY

In non-neuronal cells, clathrin has established roles in endocytosis, with clathrin-cages enclosing plasma-membrane infoldings, followed by rapid disassembly and reuse of monomers. However, in neurons, clathrin is conveyed in slow axonal transport over days to weeks, and underlying transport/targeting mechanisms, mobile cargo-structures – and even its precise presynaptic localization and physiologic role – is unclear. Combining live-imaging, photobleaching/conversion, mass-spectrometry, electron-microscopy and super-resolution, we found that unlike dendrites where clathrin-cages rapidly assemble and disassemble; in axons, clathrin and related proteins organize into stable ‘transport-packets’ that are unrelated to endocytosis and move intermittently on microtubules, generating an overall slow anterograde flow. At synapses, multiple clathrin-packets abut synaptic-vesicle (SV) clusters, and clathrin-packets also exchange between

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AUTHOR CONTRIBUTIONS:

S.R. conceived the project. A.G., R.S., and S.R. designed most of the experiments and analyzed the data. C.L., F.W., and G.C. designed and performed the DNA-PAINT experiments and data-analyses. A.G. performed the live-imaging experiments in axons and dendrites, and analyzed most of the data. Electron microscopy/analyses was performed by A.G. with assistance from D.B. and S.P. at NCMIR, led by M.H.E. Samples for mass-spectrometry were prepared by A.G.; experiments were performed by X.H. under the supervision of J.Y.; and ensuing data were analyzed by X.H. and A.G. R.S. performed the synaptic live-imaging, photo-manipulation, FKBP/FRB/FM-uptake, and live/dead-assay experiments; analyzed the ensuing data; and also helped analyze some of the axonal transport data. N.B. designed strategies to analyze the photo-manipulation experiments and performed most of the ensuing analyses, with assistance from R.S. and S.R. L.P. helped with neuronal cultures, and U.D. and L.P. designed and generated constructs. S.R. wrote the manuscript, with contributions from C.L., R.S. and A.G.

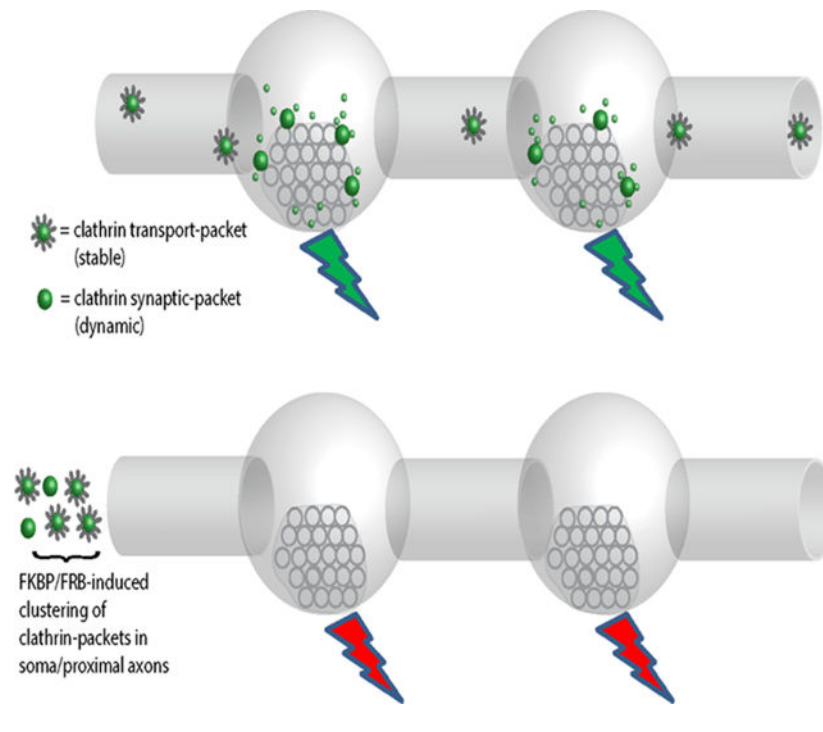
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synaptic boutons in a microtubule-dependent ‘superpool’. Within synaptic boundaries, clathrin is surprisingly dynamic, continuously exchanging between local clathrin-assemblies, and their depletion impairs SV-recycling. Our data provide a conceptual framework for understanding clathrin trafficking and presynaptic targeting that has functional implications.

In brief:

Clathrin helps maintain synaptic homeostasis, but it’s unclear how it gets to synapses. Ganguly and Sharma et al. show that clathrin assembles into stable transport-packets in axons that are trapped upon reaching synapses, where packets radially organize around synaptic-vesicle clusters. Synaptic clathrin-packets are dynamic, and their depletion impairs synaptic function.

Graphical Abstract



INTRODUCTION

The cytosolic protein clathrin is an established player in endocytosis. During this process, soluble clathrin is recruited to the inner plasma-membrane, forming clathrin-coated pits and vesicles. After vesicle internalization, the clathrin coat is rapidly removed by uncoating proteins, and the released clathrin monomers are reused for subsequent rounds of endocytosis. Clathrin-mediated endocytosis (CME) has been studied for over forty years, and a host of adapters and regulators are known. Clathrin also plays roles in sorting of intracellular membranes, although this is less studied (Jung and Haucke, 2007; Dittman and Ryan, 2009; McMahon and Boucrot, 2011; Traub and Bonifacino, 2013).

In neurons, clathrin synthesized in cell-bodies is transported into axons – enriching at presynapses – and previous *in vivo* pulse-chase radiolabeling studies have established that clathrin is conveyed in slow axonal transport, taking days to weeks in reaching axon terminals (Black et al., 1991; de Waegh and Brady, 1989; Elluru et al., 1995; Garner and Lasek, 1981; Gower and Tytell, 1987). Slow transport is a poorly defined rate-class carrying cytosolic (or soluble) and cytoskeletal proteins to the axon-tips and presynaptic terminals – unlike fast vesicular cargoes that are rapidly transported in minutes to hours [reviewed in (Maday et al., 2014; Roy, 2014, 2020)]. Although radiolabeling techniques established the overall movement of clathrin in slow transport, these methods could not visualize the movement, and many questions remain.

How can clathrin undergo organized slow axonal transport, given the ephemeral nature of cytoplasmic clathrin-assemblies? Indeed, previous studies in cultured hippocampal neurons showed rapid on/off behavior of GFP:clathrin puncta in dendrites, lasting only for seconds, consistent with CME (Blanpied et al., 2002; Rosendale et al., 2017). While clathrin may organize into longer-lasting assemblies for slow transport, no such structures have been described. How is clathrin targeted to presynapses and retained there? What is its nanoscale organization at synapses? What is the dynamic behavior of clathrin at the synapse, and can this offer insight into function? Though clathrin is involved in retrieving SVs from the presynaptic plasma-membrane (Heuser and Reese, 1973; Royle and Lagnado, 2010; Saheki and De Camilli, 2012), newer studies argue that clathrin acts downstream of endocytosis – in the regeneration of new SVs (Chanaday et al., 2019; Chanaday and Kavalali, 2018; Milosevic, 2018) – and a better understanding of the precise localization and dynamic behavior of clathrin may offer a fresh perspective.

Recent studies from us and others using cultured neurons as a model-system – to visualize and manipulate slow axonal transport – have begun to offer some answers into mechanisms (Chakrabarty et al., 2019; Ganguly et al., 2017; Scott et al., 2011; Tang et al., 2013; Twelvetrees et al., 2016). Here, we report previously unknown long-lasting axonal assemblies of clathrin and associated proteins that are unrelated to endocytosis and specialized for cargo-delivery. We also found a striking radial organization of clathrin around – but not within – SV-clusters, and a microtubule-dependent clathrin superpool that may be distinct from the known actin-dependent SV-superpool (Darcy et al., 2006). Though the transported axonal clathrin-packets are stable, surprisingly, they become dynamic upon entering – exchanging clathrin molecules with local synaptic clathrin – and depletion of clathrin-packets from boutons impairs SV-recycling. Collectively, the data advocate new models for trafficking and targeting of clathrin and related proteins, with implications for synaptic function.

RESULTS

Discrete clathrin transport-packets move intermittently in slow axonal transport

Although the overall appearance of clathrin-puncta in cultured neurons seems similar in somatodendritic and axonal compartments by immunostaining (Fig. 1A), their dynamics were different. To visualize clathrin in living neurons, we used GFP (or mCherry) tagged to the N-terminus of clathrin light chain-A (CLC) – a structurally and functionally active

fusion protein (Gaidarov et al., 1999) used in other studies (Blanpied et al., 2002; Mueller et al., 2004; Pelassa et al., 2014; Rosendale et al., 2017; Zhao and Keen, 2008). All live imaging of axons was performed between 8–10 days in vitro (DIV), and synapses were imaged between 13–21 DIV – a time when synapses are established in our cultures. As reported previously (Blanpied et al., 2002; Rosendale et al., 2017), GFP:CLC showed largely on/off dynamics in dendrites, reflecting transient recruitment of clathrin to coated-pits in kymographs (Fig. 1B; also see Supp. Movie 1). However, axonal GFP:CLC particles were long-lasting, and ~ 40% were motile; moving with an overall anterograde bias (Fig. 1C–E, also see Supp. Fig. 1A and Supp. Movie 2). Where visible, moving clathrin particles did not colocalize with transported vesicles (Supp. Fig. 1B). To distinguish moving axonal clathrin structures from dendritic clathrin-coated pits and other endocytosis-related clathrin structures, here we call them clathrin ‘transport-packets’.

Previously, we developed an assay to visualize the anterogradely-biased transport of cytosolic cargoes moving in slow transport (“intensity-center shift assay”; see Roy et al., 2012; Scott et al., 2011). Here, cultured neurons are transfected with photoactivatable (or photoconvertible) probes tagged to the cytosolic protein of interest, and a discrete region of interest (ROI) in the axon-shaft is photoactivated/photoconverted (schematic in Fig. 1F). Thereafter, a line is drawn along the long-axis of the axon including the photoactivated/photoconverted ROI, and the intensity-center (or centroid) shift of the fluorescence in the ROI is monitored over time. While untagged photoactivatable GFP (PAGFP) rapidly diffuses without any centroid-shift, cytosolic proteins disperse with a slow anterogradely-biased centroid-shift; consistent with slow transport (Chakrabarty et al., 2019; Ganguly et al., 2017; Scott et al., 2011; Tang et al., 2012; Tang et al., 2013; Twelvetrees et al., 2016). A kymograph from these experiments is shown in Fig. 1G, along with the ensuing raw and ensemble data (Fig. 1H–I). Note that PAGFP:CLC – but not soluble, untagged PAGFP – has an overall anterograde bias with a predicted average rate of ~ 0.006 $\mu\text{m/s}$ or ~ 0.5 cm/day, in line with overall rates of slow transport in radiolabeling studies (Garner and Lasek, 1981). Similar results were obtained with a photoconversion assay where we first visualized green Dendra2:CLC puncta in axons and then photoconverted them to red (Fig. 1J). Collectively, the data indicate that the intermittent movement of axonal clathrin transport-packets generates an overall anterograde bias of the clathrin population, at rates consistent with slow axonal transport.

Motility of clathrin transport-packets is independent of endocytosis

The persistence and biased movement of the transport-packets suggest that clathrin is conveyed as a unique structure that is specialized for axonal transport, with no role in endocytosis. To test this idea, we asked if inhibition of endocytosis would also affect the motility of axonal transport-packets. First, we globally suppressed endocytosis in cultured neurons using Dynasore – a small-molecule inhibitor of dynamin (Das et al., 2013; Macia et al., 2006). We also specifically inhibited clathrin-dependent endocytosis with CHC-T7-Hub, a clathrin heavy-chain dominant-negative mutant known to disrupt clathrin-dependent endocytosis (Bennett et al., 2001). As reported previously in non-neuronal cells, the CHC-T7-Hub mutant attenuated receptor-mediated endocytosis in cultured hippocampal neurons (Supp. Fig. 2A–B). The on/off dynamics of GFP:CLC in dendrites was essentially abolished

in neurons treated with Dynasore (Fig. 2A), or transfected with the CHC-T7-Hub mutant (Supp. Fig. 2C–D). Interestingly however, neither Dynasore, nor the CHC-T7-Hub mutant had any effect on the mobility of axonal clathrin transport-packets (Fig. 2B–D). Since the motility of individual clathrin transport-packets generate the overall anterogradely-biased flow of the axonal clathrin population (see Fig. 1H), a prediction is that endocytosis-inhibition would not interfere with the slow transport of clathrin. Indeed, neither Dynasore nor the CHC-T7-Hub mutant had any effect on the biased transit of PAGFP:CLC in axons (Fig. 2E). Previous studies suggest that the slow transport of synapsin and dynein is microtubule-dependent (Scott et al., 2011; Twelvetrees et al., 2016). Low-levels of the microtubule-disrupting drug Nocodazole – that blocks vesicle transport, see Supp. Fig. 2F – also inhibited movement of clathrin transport-packets (Fig. 2F). Taken together, the data indicate that axonal clathrin transport-packets are unique conveyance structures; unrelated to endocytosis.

Structure of axonal clathrin transport-packets

What do the transport-packets look like? To visualize the ultrastructure of axonal clathrin, we tagged clathrin to Apex – an engineered peroxidase that acts as an electron-microcopy (EM) tag by catalyzing the polymerization and local deposition of diaminobenzidine (DAB); subsequently recruiting electron-dense osmium for EM contrast (Martell et al., 2012). Cultured neurons were transfected with Apex-GFP:CLC, fixed and incubated with DAB and H₂O₂ (see schematic in Fig. 3A and methods). Transfected neurons were visualized by GFP fluorescence and DAB-staining; dendrites and axons were identified by morphology; and the same neurons and processes were visualized by EM tomography (see Supp. Fig. 3 and “Methods”). As expected, coated pits and coated-vesicles were seen in soma and dendrites (Supp. Fig. 2B). However, axons contained intact clathrin-coated structures that were inside the axon-shaft, with no association with the axonal plasma membrane, as seen by EM-tomography (Fig. 3B–B’; also see 3-D view in Supp. Movie 3). The diameter of these clathrin-structures as determined by EM was ~ 50nm and ~125nm (inner spherical core and outer diameter with spokes respectively); consistent with previous cryo-EM studies of clathrin coated-vesicles (Heymann et al., 2013).

We also used DNA-PAINT, a single-molecule based super-resolution microscopy (Jungmann et al., 2014) to visualize the nanoscale organization of endogenous clathrin along axons. Axonal boundaries were determined by visualizing β 2-spectrin, a component of the submembrane periodic axonal scaffold [reviewed in (Leterrier et al., 2017)]. Unlike the tedious Apex EM-imaging process involving light/EM correlation of transfected axons, DNA-PAINT allowed us to visualize and characterize a large number (>500) of axonal clathrin structures. From 3D multicolor DNA-PAINT images of clathrin and β 2-spectrin, we generated 3D-renderings for the segmentation and measurement of axonal clathrin particles; some of which are likely to be mobile transport-packets (Fig. 3C). The average equivalent diameter of clathrin heavy-chain labeled packets was 43.3 \pm 0.5 nm (mean \pm SEM, 1294 packets) – consistent with the core-size of the structures seen in our EM data – and the average density was 7.6 \pm 0.5 packets/ μ m of axon (82 rendered axonal segments). Interestingly, most clathrin packets seen by super-resolution imaging were close to the β 2-spectrin sheath, suggesting associations with the submembrane actin/spectrin scaffold

(see cross-sections of axons in Fig. 3C and Supp. Movie 4). Given that the majority of axonal clathrin particles are immobile at any given time, we speculate that this may reflect anchoring of clathrin assemblies to the periodic actin/spectrin lattice.

Composition of axonal clathrin transport-packets

To define the composition of axonal clathrin, we immunoprecipitated (IP) clathrin from mouse sciatic axon fractions, followed by MUDPIT-MS [MultiDimensional Protein Identification Technology-Mass Spectrometry, (Liao et al., 2008; Yates et al., 2009)]; see protocol in Supp. Fig. 4A. A diverse group of cytosolic proteins were identified (Fig. 4A; see Supp. Table 1 for full list – also submitted in PRIDE database, PXD005011), and as expected, clathrin was one of the most abundant protein (Fig. 4B). 115 proteins were identified, and many were cytosolic proteins that are known to reside in synapses (Wilhelm et al., 2014). Network analyses showed that the axonal clathrin-interactome contained many cytoskeleton-associated proteins such as actin isoforms and myosins (Fig. 4A, dashed circle). Actin and clathrin have established anatomic and functional links in non-neuronal cells [reviewed in (Kaksonen et al., 2006)]. Previously, using a probe for filamentous actin (GFP tagged to the calponin homology domain of utrophin, GFP:Utr-CH), we found that axon-shafts have focal actin “hotspots” every ~ 3–4 μ m along its length, where actin assembles and disassembles continuously, and that these foci serve as a nidus for nucleation of actin filaments that elongate bidirectionally along the axon-shaft [“actin trails”, see (Ganguly et al., 2015)]. Interestingly, simultaneous imaging of GFP:Utr-CH and GFP:CLC showed that there was a striking colocalization of actin hotspots and stationary clathrin in axons, though the actin foci were much more short-lived as expected (Fig. 4C; Supp. Fig. 4B).

Network analyses also revealed that axonal clathrin was associated with proteins known to disassemble clathrin coated-pits (such as GAK, Hsc70, synaptojanin), as well as those linked to clathrin pit-formation (dynammin, AP-complex); see Fig. 4A. The presence of both assembly and disassembly proteins in MS-datasets is intriguing, as the transported clathrin-structure is clearly discrete. To test if candidate assembly/dis-assembly proteins from the M/S dataset are indeed co-transported with clathrin, we visualized the transport of mCherry:CLC with GFP-fusions labeling actin, GAK, AP180 and dynammin, using simultaneous two-color live imaging. A significant fraction of GAK was also co-transported with clathrin (Fig. 4D). Like clathrin, the dynamics of GAK in dendrites was also very different from axons. As shown in the kymograph in Supp. Fig. 4C, foci of GFP:GAK typically coincided with clathrin disassembly in dendrites, very different from the mixture of stationary and motile GAK particles seen in axons. Quantification showed that significant amounts of GAK, AP180 and dynammin were co-transported with clathrin (Fig. 4E). In previous proteomic studies, we identified Hsc70 as a regulator of the slow axonal transport of synapsin (Ganguly et al., 2017). However pharmacologic or genetic inhibition of Hsc70 had no effect on the axonal GFP:CLC motility (Supp Fig. 4D), suggesting that the mechanistic basis for the transport of the two slow-component cargoes – clathrin and synapsin – is different. Indeed, synapsin has an anterogradely biased diffusion-like flow in axons (Scott et al., 2011; Tang et al., 2013), while clathrin is conveyed as discrete, stable packets.

Nanoscale organization of synaptic clathrin

Clathrin is strongly enriched at presynaptic boutons and terminals, though its precise function at this locale has been controversial in recent years (Chanaday et al., 2019; Milosevic, 2018; Watanabe et al., 2013). By light microscopy, we noticed that the presynaptic localization of clathrin was not homogenous – like the classical vesicular or cytosolic synaptic markers synaptophysin or synapsin – but appeared as multiple puncta within a bouton. To pinpoint the spatial relationship of *endogenous* presynaptic clathrin to synaptic-vesicle clusters, we first visualized synapses and adjacent axons at nanometer resolution in 3-D, with two-color DNA-PAINT; using antibodies to clathrin and synapsin (latter to label SV-clusters, see methods for details). In the cropped axon and *en-passant* bouton shown in Fig. 5A (entire image in Supp. Fig. 5A), note that multiple clathrin particles (magenta) are abutting the SV-cluster (green). 3-D views (Fig. 5B and Supp. Movie 4) unequivocally show that clathrin particles are circumferentially organized around SVs, and rarely seen deep within the SV-cluster. Segmented rendering of the super-resolution data showed that each presynapse contains ~ 4–12 clathrin particles (Fig. 5C) that were on average ~ 43 nm in equivalent diameter (Fig. 5D), consistent with the size of clathrin assemblies seen in axon-shafts. The morphologic similarities between the axonal and presynaptic clathrin particles suggest that they are related to transport-packets, and hereafter we refer to them as synaptic clathrin-packets. The large number of synaptic clathrin-packets visualized by DNA-PAINT (>1200 packets from >100 synapses) also allowed us to explore their correlation with SV-clusters. The total volume of all clathrin-packets at a given synapse correlated with volume of the SV-cluster (Fig. 5E), suggesting putative links between clathrin-levels and SV-pools. Taken together, the data suggest that motile axonal transport-packets are targeted to presynapses, where they radially organize around SVs.

Though we were not able to confidently visualize the Apex-EM signal at synapses due to high background, we analyzed immuno-gold EM datasets collected by Dr. Tao-Cheng at NINDS/NIH (also see Tao-Cheng, 2020). As shown in the representative images (Fig. 5F), immuno-gold particles were mostly seen around SV-clusters, and some gold particles were associated with vesicular structures resembling endosomes and cisternae (asterisk in Fig. 5F2). Coated vesicles are rarely seen at synapses, as reported by previous EM studies (Heuser and Reese, 1973; Kononenko et al., 2014; Petralia et al., 2003; Tao-Cheng, 2020). Taken together, the super-resolution and immuno-EM data, along with previous studies, indicate that endogenous synaptic clathrin localizes at the periphery of SV-clusters, and the majority of synaptic clathrin is not in the form of coated-vesicles.

Clathrin-packets are captured and immobilized at synapses

To explore the dynamics of synaptic clathrin, we first visualized GFP:CLC at synapses with relatively high temporal resolution (5 frames/sec). Motile clathrin particles rapidly moved between boutons (average velocity 0.54 $\mu\text{m/s}$) and were often trapped at synapses (kymographs in Fig. 6A and Supp. Fig. 6A). A “pass/trap analyses” (see Supp. Fig. 6B and “Methods”) indicated that ~60% of the moving particles were retained at boutons for at least a minute (129 particles analyzed from 19 neurons, 3 separate cultures). Within a bouton, clathrin particles moved back and forth – seemingly constrained by the synaptic boundary – though they could exit (Fig. 6A’). Single synaptic GFP:CLC particles could only be tracked

for a few seconds, suggesting instability (see photoconversion data later). Next, we used FRAP to bleach GFP:CLC in a single bouton and visualized fluorescence-recovery over time – reflecting the entry of unbleached fluorescent molecules from adjacent boutons/axons into the bleached-zone. As shown in Fig. 6B, there was a slow recovery of fluorescence over several minutes, and particles were frequently seen to enter the bleached bouton (best seen in Supp. Movie 5). Interestingly, FRAP of GFP:CLC fluorescence at boutons was qualitatively very different from GFP:synapsin. While recovery of clathrin was largely due to entry of discrete fluorescent structures – and was somewhat stochastic – synapsin FRAP was gradual and diffuse (Fig. 6C and Supp. Fig. 6C). Also, clathrin mobility in presynaptic boutons decreased over time in our cultures (Supp. Fig. 6D), likely due to synaptic maturation, reminiscent of the on/off behavior of GFP:CLC in dendrites, that also decreases with time (see Blanpied et al., 2002).

Fluorescence recovery of a freely-diffusing molecule is expected to be complete (see FRAP of soluble, untagged GFP in boutons; Fig. 6D - inset), and incomplete recovery reflects a bound, immobile pool that was unable to enter the bleached zone (Lippincott-Schwartz et al., 2003). While FRAP of both clathrin and synapsin was incomplete, clathrin had a much larger immobile fraction (Fig. 6D–E and Supp. Fig. 6E). A previous study in *C-elegans* found that endophilin – a soluble protein involved in endocytosis – showed diffusion-like mobility between synapses that was blocked when SV-recycling was perturbed (Bai et al., 2010); suggesting that the trafficking endophilin pool was generated locally after SV-recycling. Though the diffusion-like kinetics of endophilin is very different from the particle-movement of clathrin, it is possible that the inter-synaptic movement of clathrin is also dependent on endocytosis, and that the clathrin-packets are locally generated at the synapse. To test this, we performed clathrin FRAP experiments before and after adding Dynasore. Dynasore did not attenuate FRAP of clathrin; and unexpectedly, there was an increase in clathrin motility (Fig. 6F). One possibility is that blocking endocytosis might decrease the usage of clathrin-packets at synapses, allowing more packets to shuttle between synapses (perhaps also increasing their synaptic capture). Indeed, after Dynasore-treatment, there was an increase in the number of clathrin-packets exchanging between boutons – i.e. exported from one synapse and captured at another (Fig. 6G). Collectively, the data suggest that the “superpool clathrin-packets” are related to trafficking, and not generated at synapses following endocytosis. The increase in the synaptic capture after Dynasore also suggests that synaptic targeting of clathrin may be independent of its function at the synapse.

The morphology and dynamics of synaptic clathrin puncta suggest that axonal transport and subsequent presynaptic “trapping” of a discrete number of transport-packets may be the basis of clathrin enrichment at boutons. To further explore morphologic similarities between the transported clathrin and the clathrin localized at synapses, we asked if the total fluorescence of clathrin at individual boutons was an aggregate of integer-multiples of axonal puncta/transport-packets. Overlaid histogram-distributions of GFP:CLC fluorescence intensities in axons (green) and presynapses (red) are shown in Fig. 6H (left). Note that while GFP:CLC axonal intensities occupy a single-peak, synaptic GFP:CLC intensities are distributed as multiple peaks of higher intensities. Fig. 6H (middle) shows predicted fluorescence peaks of synaptic clathrin if they were integer-multiples of the axonal intensity peak (assumed as ‘x1’). From this analyses, single boutons are predicted to contain ~ 4–8+

clathrin-packets (Fig. 6H, right). A similar distribution of axonal and synaptic fluorescence was also seen for endogenous clathrin (Supp. Fig. 6F). Collectively, the data are consistent with a model where synaptic enrichment of clathrin is largely due to clathrin-packets shipped by slow axonal transport.

Dynamics of synaptic clathrin-packets

Next, we used photoconvertible clathrin to determine the fate of synaptic clathrin. Neurons were transfected with Dendra2:CLC (green to red photoconversion), and synaptic clathrin was identified by constitutive green fluorescence (see “Methods” for details). Dendra2:CLC in a single bouton was photoconverted to red, and both green (stable) and red (photoconverted) fluorescence was monitored over time. Photoconverted clathrin particles moved out of the synapse into the flanking axons, and were often trapped at adjacent boutons (Fig. 7A, also see Supp. Movie 6). Pass/trap analysis showed that ~67% of moving particles were trapped (see Supp. Fig. 6B and “Methods for details; 46 particles analyzed from 14 neurons; 3 separate cultures). The overall dispersion of clathrin from photoconverted boutons – determined by decay of red fluorescence over time – is shown in Fig. 7B. Previous studies have described a mobile vesicle superpool spanning multiple synaptic boutons and flanking axons, composed of vesicles that are available to synapses during stimulation (Darcy et al., 2006; Krueger et al., 2003; Staras et al., 2010); and pharmacologic experiments suggest that the SV-superpool is actin-dependent (Darcy et al., 2006; Ratnayaka et al., 2011). Though the kinetics of the clathrin particles in and around synapses resemble the vesicle-superpool, interestingly, the peri-synaptic motility of photoconverted Dendra2:CLC particles was microtubule – and not actin – dependent (Fig. 7C–D).

To explore the dynamics of individual synaptic-packets, we photoconverted a single Dendra-2:CLC packet within a bouton and visualized both stable (green) and photoconverted (red) fluorescence by live-imaging. Given the stability of the axonal transport-packets, we expected that the fluorescence would be stable in the photoconverted synaptic packet as well. Surprisingly however, the photoconverted synaptic packet lost fluorescence over tens of seconds, while the other clathrin-packets within the same bouton – identified by the stable green fluorescence – gained fluorescence over the same time-frame (Fig. 7E and Supp. Movie 7, more examples in Supp. Fig. 7A, A’). Clathrin exchange between synaptic packets was not seen in boutons from neurons fixed with paraformaldehyde (Supp. Fig. 7B). To quantify the aggregate data from these experiments, we compared the change of fluorescence in the photoconverted packet in a bouton to that of the non-converted (“dark”) packets within the same bouton (see basic principle in Fig. 7F schematic). If there was exchange, fluorescence would be lost from the photoconverted packet, and there would be a gain of fluorescence in the “dark” regions. As shown in the graphs in Fig. 7F, fluorescence was lost from photoconverted packets in living neurons (left red bar) and gained in the non-photoconverted dark regions (left black bar). However, in fixed boutons, fluorescence was not lost from the photoconverted packet (right red bar, Fig. 7F), and there was little change of fluorescence in the non-photoconverted dark regions of the synapse (right black bar, Fig. 7F). Occasionally in axons, there was a mechanical dispersion of the axonal fluorescence due to other moving axonal packets that happened to

cross the field of view (see Supp. Fig. 7C for an example). A side by side comparison of synaptic and axonal Dendra2:CLC photoactivation is shown in Supp. Fig. 7D. Collectively, the data suggest that though synaptic clathrin packets are derived from axonal packets, the dynamic behavior of the packets changes upon arrival at the synapse.

Physiologic role of clathrin-packets

Do the clathrin transport/synaptic-packets have a physiologic role at synapses? To address this, we designed a system to sequester clathrin-packets in the neuronal soma – depleting them from synapses – and then examined SV-recycling in the clathrin-depleted boutons using FM4–64; a red styryl dye that is selectively endocytosed into rapidly recycling SVs and has been widely used to evaluate synaptic function (Gaffield and Betz, 2006). Though previous studies have used shRNAs to deplete clathrin over several days, clathrin is required for Golgi-formation (Radulescu et al., 2007) and essential for survival (Bazinet et al., 1993). Thus we sought a more acute knockdown of clathrin, and also a method that depletes clathrin-packets from synapses. Towards this, we used the FKBP-Rapalog-FRB heterodimerization system, where proteins fused to FKBP or FRB dimerize upon adding Rapalog (Belshaw et al., 1996; Bentley et al., 2015; Kapitein et al., 2010; Robinson et al., 2010). We tagged FRB to GFP:CLC (FRB-GFP:CLC), and FKBP to the dynein-binding fragment of BicD2 (BicD2-FKBP +/- Tdtomato tag), a dynein cargo-adaptor that moves cargoes towards the minus-end of microtubules (Dienstbier and Li, 2009).

In our system, Rapalog would induce dimerization of GFP:CLC labeled clathrin-packets and BicD2, transporting them towards the minus-end of axonal microtubules and into the neuronal soma – thus depleting clathrin-packets from boutons (see schematic in Fig. 8A). Fig. 8B shows examples of neurons transfected with FRB-GFP:CLC and Tdt:BicD2-FKBP, with or without Rapalog. Note sequestration of GFP:CLC in neuronal cell-bodies and its depletion from synapses after incubation with Rapalog (Fig. 8C – quantified in Fig. 8D). Importantly, almost all of the transfected FRB-GFP:CLC co-assembles with endogenous axonal clathrin (Supp. Fig. 8A–B); thus sequestering FRB-GFP:CLC leads to a depletion of *all* synaptic clathrin in these experiments, including endogenous clathrin (see Fig. 8E). Neurons appeared healthy with normal axonal and dendritic arborizations after FKBP/FRB/Rapalog mediated clathrin sequestration for up-to 10h, and a live/dead assay (Liljebäck et al., 2019) showed no toxicity (Supp. Fig. 8C).

Next we evaluated SV-recycling in the clathrin-depleted boutons using FM-uptake protocols that specifically label the recycling pool [Cousin et al., 2018; see Fig. 8F and “Methods” – for discussion of various synaptic pools, see Denker and Rizzoli, 2010]. Incubation with Dynasore led to ~75% decrease in FM-uptake (Supp. Fig. 8D); and as shown in Fig. 8G, there was also a significant reduction in FM-dye uptake in the Rapalog-treated neurons (quantified in Fig. 8H – similar changes were seen at 37°C, Supp. Fig. 8E). Interestingly, though Rapalog treatment for 10h led to a greater attenuation of synaptic clathrin when compared to the 5h Rapalog-treatment (see Figs 8D–E and 8G – left graph), the extent of FM attenuation was similar at both 5h and 10h timepoint, suggesting that there may be a finite functional pool of synaptic clathrin, and the remaining pool might

help maintain homeostatic protein-levels at this locale. Taken together, the data suggest that clathrin-packets at the synapse are involved in maintaining SV-recycling.

DISCUSSION

Our main findings are: i) clathrin and related proteins are conveyed in axons as stable coated structures (transport-packets) – moving intermittently on microtubules with an overall slow anterograde bias; ii) at synapses, most clathrin assemblies (synaptic-packets) localize to the periphery of SV-clusters, and despite morphologic similarities with transport-packets, most synaptic-packets lack a conventional coat; iii) mobile clathrin-packets shuttle between synapses in a microtubule-dependent superpool, and a significant fraction (~ 60%) of the moving clathrin is captured at boutons; iv) though axonal clathrin-packets are stable, synaptic clathrin-packets can exchange with other packets within a bouton; and v) depletion of synaptic clathrin-packets impairs SV-recycling.

A model for trafficking and presynaptic targeting of clathrin

The most straightforward model emerging from our data is that clathrin and related proteins are synthesized in the neuronal soma, where they are packaged into stable coated structures that are dedicated for transport – with no role in endocytosis. Transport-packets move intermittently on microtubules, and are conveyed to synapses with an overall anterograde bias in slow axonal transport. Upon reaching the synapse, the conventional coat on the packets is disrupted – perhaps due to the high concentration of uncoating proteins like Hsc70 at this locale (Ganguly et al., 2017) – allowing clathrin to dynamically exchange with other local packets. Thus, at synapses, clathrin molecules would either be associated with a synaptic-packet, or exchange between local packets. This metastable state would provide a labile local pool of clathrin that could be rapidly recruited for synaptic function, as needed. In this hypothetical model, clathrin would undergo a seamless transition from transport to function upon reaching the synapse, simply by dynamically shedding the coat on the transport-carrier.

Despite the dynamic exchange, our experiments strongly suggest that the synaptic clathrin is not freely diffusible as commonly believed. For instance, in photoconversion experiments (Fig. 7A), we only saw discrete clathrin particles leave boutons, and incubation with Nocodazole – that stalled the mobility of the synaptic clathrin-packets – did not lead to a loss of clathrin from the stalled structures over time (Fig. 7C), which would be predicted if this clathrin was freely diffusible. However, the precise structures on which the synaptic clathrin localizes is still unclear, though immuno-EM data suggest that some of the clathrin may be on cisternae/endosomes (Fig. 5F). Our M/S data and two-color live imaging also suggest that many proteins related to endocytosis are co-transported with clathrin, so once the transport-packets reach the synapse, perhaps they too would be available for performing their functions. Thus, conceptually, our model may have broad consequences for neuronal form and function.

Synapses have multiple integers of clathrin transport-packets (Fig. 6G), which is also consistent with our working model where clathrin trafficking and targeting are interconnected events. However, EM suggests that while axonal clathrin-packets are coated,

most of the synaptic clathrin lacks a conventional coat. Previous studies have shown that clathrin coated-vesicles are infrequent in resting hippocampal synapses – for example in a recent detailed study, only ~5–10 coated-vesicles were seen per 100 presynaptic profiles (Tao-Cheng, 2020) – but it is clear that clathrin is abundant at synapses, obvious by routine immunostaining. Our live-imaging, super-resolution and EM data provide some clarity on this issue, showing that most of the synaptic clathrin: 1) is organized into discrete structures, and not as free monomers; and 2) does not resemble conventional clathrin-coated vesicles. Another prediction of our model is that during transport, the stable clathrin on axonal transport-packets would not be functionally available, which might be a way of preventing unwanted endocytosis along the axon shaft.

Clathrin Transport-Packets – a Carrier for Slow Axonal Transport

Though the phenomenon of cytosolic slow transport is established, the nature of transported cargoes has been puzzling. In previous imaging studies of other slow-component cargoes (like synapsin and CamKII), we saw transient mobile structures – lasting for a few seconds at best – that appeared to contribute to the overall anterograde movement (Roy, 2013; Scott et al., 2011), but the exact cargo-structure remained unclear. The discrete nature of axonal clathrin enabled us to examine the cargo-structure at a nanoscale level by 3D-EM and super-resolution, and clathrin is probably the first cytosolic cargo to be characterized in detail. Based on the overlapping of radiolabeled “peaks” in axons, previous studies hypothesized that common mechanisms were involved in slow transport [reviewed in (Roy, 2014, 2020)]. However, the synaptic dynamics of two slow transport cargoes clathrin and synapsin is very different (Fig. 6C), and an emerging theme in slow transport is the diversity in the nature and mechanisms of movement (Roy, 2020).

The structure of an axonal transport-packet resembles clathrin coated-vesicles. However, coated-vesicles are thought to be transient intermediates that deliver endocytosed vesicles – and their contents – either to Golgi/endosomes, or back to the plasma-membrane [reviewed in (Peraan et al., 2020)], while the transport-packets are clearly long-lasting and being delivered to distal synapses. Though previous studies in non-neuronal cells have described motile coated intermediates in the cytoplasm (Keyel et al., 2004; Puertollano et al., 2003; Rappoport and Simon, 2003; Rappoport et al., 2003a; Rappoport et al., 2003b), these seem different from the axonal transport-packets. First, the intermediates are described as large, pleomorphic tubulovesicular aggregates (Polishchuk et al., 2006; Puertollano et al., 2003); different from the ~40–50 nm uniform-sized structures we see in the axon (Fig. 3). Also, the clathrin in these moving intermediates is reported to “cycle on and off” – eventually fusing with early or late endosomes (Puertollano et al., 2003) – which is also different from the stable structures we see in axons. However, it is possible that the transported clathrin-packet is a stable form of specialized coated-vesicle dedicated to slow transport.

Alternatively, the transported structure might represent clathrin assemblies that are not around vesicles. Indeed, the lumens of clathrin transport-packets do not appear electron-dense (see Fig. 3B”); though only cryo-EM would be conclusive. Interestingly, “clathrin baskets” – clathrin cages with no vesicles inside – are routinely isolated from biochemical fractionations of bovine brains (Nandi et al., 1982), and in fact, comprise ~ 80% of

the clathrin structures isolated in these preparations (Heymann et al., 2013). Clathrin baskets are thought to be artefacts, assembled in the homogenate during biochemical isolation (Heymann et al., 2013). However, given the long-lasting coated transport-packets we describe here, and the large combined relative-volumes that axons must occupy in brain homogenate preparations, it is conceivable that clathrin-baskets are actually axonal transport-packets. One caveat in our experiments is that our Apex-EM experiments cannot distinguish between transported and stationary clathrin in the axon.

Mechanistic implications of Clathrin Trafficking and Targeting

The movement of clathrin between presynaptic boutons resembles SV-superpools, that are thought to be an extension of local SV-pools (Staras et al., 2010). However, while the SV-superpool is dependent on actin dynamics, the “clathrin superpool” depends on microtubules (Fig. 7D) and may be a distinct superpool conveying endocytic (and perhaps other slow-component) proteins. Interestingly, the mobility of clathrin particles is also different from the soluble diffusive motion of endophilin (Bai et al., 2010), so it seems likely that there are many components to the superpool. Substantial evidence from our live-imaging, photobleaching and photoconversion experiments support the view that trafficking clathrin-packets are captured at synapses (Figs. 6–7), and their depletion causes functional deficits (Fig. 8); thus it seems reasonable that the trafficking pool is critical for synaptic function. We posit that multiple synaptic clathrin-packets that may be “poised” to provide a pool of clathrin that would be locally available in the vicinity of SVs; thus speeding up SV-recycling. Previous studies suggest that clathrin may be locally translated at axons and synapses (Biever et al., 2020; Shigeoka et al., 2016), and our experiments do not exclude a possible role for local translation. The function of clathrin at synapses has been controversial in recent years, and clathrin has been proposed to play roles in SV biogenesis, particularly at physiologic temperatures (Delvendahl et al., 2016; Soykan et al., 2017; Watanabe et al., 2013). In that context, our key experiments were done at ~ 37°C, though our experiments cannot directly distinguish between the different models. Finally, the behavior of synaptic clathrin resembles the movement of soluble molecules within liquid condensates and may be conceptually related to the contemporary idea that SV-clusters have fluid-like properties (Milovanovic et al., 2018). Interestingly, recent studies in non-neuronal cells indicate that CME also relies on the formation of liquid condensates (Day et al., 2021; Witkowska and Haucke, 2021), and future experiments focused on these issues might bring more clarity.

STAR METHODS

RESOURCE AVAILABILITY

Lead contact—Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Subhojit Roy (sroy@ucsd.edu).

Materials availability statement—Plasmids generated in this study are available from the Lead Contact without restriction.

Data and code availability

- The MUDPIT-MS data are available in the supplemental information. Microscopy data reported in this paper will be shared by the lead contact upon request.
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Animals—CD1 pups obtained from Charles River Laboratories (Cat#022-CD1) were used for preparing primary hippocampal cultures. Wistar rats were obtained from Janvier labs (France).

Primary cell cultures—For most experiments, hippocampal cultures were obtained from brains of postnatal (P0–P1) CD-1 mice and plated on MatTek glass-bottom dishes as described previously in detailed published protocols (Ganguly and Roy, 2014; Roy et al., 2012), in accordance with University of California guidelines. Briefly, MatTek dishes were coated with 100 μ l of 1 mg/ml poly-d-lysine for 2 h at RT, washed thrice with ddH₂O, and air dried before plating. Hippocampi from P0–P1 pups were dissected in ice-cold dissection buffer (HBSS, 4.44 mM d-glucose, and 6.98 mM HEPES) and incubated in 0.25% Trypsin–EDTA at 37°C for 15 min. Following this, neurons were dissociated in plating media (10% FBS and 90% Neurobasal/B27; Life Technologies) by trituration. Neurons were plated at a density of 50,000 cells/100 μ l (for FM4–64 experiments and clathrin imaging at *en-passant* boutons) and at 25,000 cells/100 μ l (for all other experiments) of plating media. Neurons were maintained in Neurobasal/B27 media (supplemented with 2% B27 and 1% GlutaMAX) in an incubator at 37°C and 5% CO₂ for 7–9 d before transfection. All the experiments on neuronal cultures were performed between 8–21 DIV. E18 rat hippocampal cultures were used for the DNA-PAINT experiments following standard protocols, and guidelines established by the European Animal Care and Use Committee (86/609/CEE) and local ethics committee (agreement D13–055-8) were followed.

METHOD DETAILS

DNA constructs, pharmacologic agents, neuronal cultures, transfections, and immunocytochemistry—The GFP:CLC (Gaidarov et al., 1999), pUC57-APEX (Martell et al., 2012) and Dendra2 constructs were obtained from Addgene [from the laboratories of Drs. James H. Keen (Thomas Jefferson University, PA), Alice Ting (Stanford University, CA) and Michael Davidson (Florida State University, Tallahassee, FL), respectively]. The PAGFP:CLC construct was subcloned from the PAGFP backbone (Patterson et al., 2002), a gift from Dr. Jennifer Lippincott-Schwartz (Janelia Farms Research Campus, Ashburn VA) by standard cloning. CHC-T7-Hub and T7-empty backbone (Liu et al., 1998) was a gift from Dr. Frances Brodsky (UCL, London, UK). The GFP:Utr-CH (Burkel et al., 2007), and synaptophysin:mCherry constructs (Gerrow et al., 2006) were gifts from Drs. William Bement (University of Wisconsin, Madison, WI) and Leon Lagnado (University of Sussex, Sussex, England, UK) respectively. pBa-Flag-BicD2-FKBP and pBa-Flag-tdTM-

BicD2-FKBP constructs were from Gary Banker (Oregon Health & Science University, Portland, OR). FRB-GFP:CLC was generated by inserting GFP:CLC in pFRB-3myc-Rab5a plasmid by replacing Rab5a (Bentley et al., 2015). GFP:synapsin-1a (Gitler et al., 2004) was a gift from George Augustine (Nanyang University, Singapore). EGFP-AP180 (Bushlin et al., 2008), GAK-GFP (Massol et al., 2006) and pEGFP-C1-Dynamin (Song et al., 2004) were kindly gifted by Drs. Pamela Yao (National Institute on Aging-National Institutes of Health, Baltimore, Maryland), Tomas Kirchhausen Harvard Medical School, Boston, MA and Sandra Schmid (University of Texas Southwestern Medical Center, Dallas, Texas) respectively. Dynasore, Latrunculin A (Life Technologies) and Nocodazole were dissolved in DMSO and used at a final concentration of 80 μ M, 100 nM and 10 μ g/ml respectively. Swinholide A and Rapamycin analogue A/C heterodimerizer (Takara) were dissolved in ethanol and used at a final concentration of 100 nM respectively. The stocks of FM4-64 (Invitrogen), Advasep-7, Live-or-Dye NucFix™ Red (Biotium) were prepared in DMSO and used at a final concentration of 10 μ M, 1 mM and 0.1x in PBS (manufacturers' protocol) respectively. The glutamate receptor antagonists 6-cyano-7-nitroquinoxaline-2,3-dione (CNQX, TOCRIS bioscience), and D,L-2-amino-5-phosphonovaleric acid (AP5, TOCRIS bioscience) were used at 10 μ M and 50 μ M concentration respectively. Unless otherwise noted, all drugs were purchased from Millipore-Sigma and added to Hibernate E media (see below) prior to imaging. The Hsc70-D10N and control Hsc70 constructs were from N. Lamarche-Vane (McGill University, Montreal, Canada). The Hsc70 inhibitor VER155008 (Cat#3803; Tocris Biosciences) was dissolved in DMSO and used at a working concentration of 100 μ M.

Neurons were transfected with the indicated fluorescent constructs (0.8 μ g DNA for clathrin constructs, 1.2 μ g of all others) at 7–9 DIV with either Lipofectamine 2000 (Life Technologies) or Calcium phosphate transfection kit (Promega) (Sharma et al., 2019). Live imaging of axons was performed between 8–10 DIV, and boutons were imaged between 13–21 DIV. As dictated by the experiment, neurons were co-transfected with soluble-GFP/soluble - mCherry/synaptophysin:DsRed to identify axons/dendrites/boutons. GFP:CLC transfected neurons selected for imaging had punctate clathrin appearance in dendrites, and axon regions imaged were more than 150 μ m from the neuronal cell body. Boutons were selected >150 μ m away from the soma on the basis of varicosities in the intensity along the axon and other morphological features typical of synapses (Wang et al., 2014).

For immunostaining, neurons were fixed in 4% paraformaldehyde/sucrose solution in 1x PBS for 10 min at room temperature, followed by extraction in 0.2% Triton X-100 for 10 min, and blocking in 1% bovine serum albumin/5% FBS for 2h at room temperature. To label endogenous clathrin, neurons were incubated overnight at 4°C with rabbit polyclonal antibody against clathrin heavy chain at 1:500 dilution. After washing the primary antibody with 1x PBS, neurons were blocked again for 30 min at room temperature, and incubated in goat-rabbit Alexa Fluor 594 (1:500) for 1h. for some experiments, GFP:CLC transfected neurons was stained with primary chicken polyclonal anti-GFP antibody (1:500) and secondary Alexa Fluor 488 (1:500). All primary and secondary antibodies were purchased from Abcam.

Live imaging, photobleaching, photoactivation, and photoconversion

experiments—Most single-color live imaging experiments for axonal trafficking were performed on an Olympus IX81 inverted motorized epifluorescence microscope equipped with a CoolSNAP *HQ2* camera (Photometrics). Two-color live-imaging was done either by simultaneous imaging of green/red channels using the Dual Cam imaging device (Photometrics) attached to two CoolSNAP *HQ2* cameras (Olympus IX81), or by rapid sequential excitation of green/red LEDs (SPECTRA-X) on an inverted epifluorescence microscope (Eclipse Ti-E; Nikon) equipped with CFI Plan Apochromat VC 100× oil (NA 1.40; Nikon) objective, and an electron-multiplying charge-coupled device camera (QuantEM:512SC; Photometrics). A multi-band-pass filter (Chroma Technology Corp.) was inserted into the emission light path, and GFP/RFP images were obtained with precise subpixel registration. Before live-imaging, neurons were transferred to Hibernate media (Brainbits), supplemented with 2% B27, 2mM Glutamax, 0.4% D-glucose, 37.5 mM NaCl (Ganguly et al., 2017; Roy et al., 2012; Scott et al., 2011), and maintained at 35.5°C to 37°C for the duration of the experiments (Precision Control, Weatherstation or a heated stage chamber, model STEV; World Precision Instrument, Inc.). Axons were identified by morphology, and only neurons with unambiguous morphology were selected for imaging (Roy et al., 2012; Scott et al., 2011). For experiments in Fig. 1, GFP:CLC, neurons were either imaged every 0.7 s for several minutes, or PAGFP:CLC was photoactivated with 405 nm LED light for 1 s, and imaged every 1.2 s. Dendra2:CLC was photoconverted by 405 nm LED light for 3 s, and imaged every 1.2 s. Synaptophysin:GFP was imaged using the stream acquisition function (MetaMorph) at 5 frames/s (with no time interval between images). For imaging *en passant* boutons, 13–21 DIV neurons were co-transfected with synaptophysin:dsRed/sol-mCherry and GFP:CLC/GFP:Synapsin, and boutons were identified based on size of the synaptic vesicle cluster and morphology (Wang et al., 2014). Boutons along the primary axon, >150 μm from the soma and <7μm apart from the each other were selected for imaging. Additionally, for the GFP:CLC live-imaging of *en-passant* boutons and intervening axonal-segments, boutons expressing >2 clathrin packets were selected and imaged at every 0.2s (rapid imaging) for several minutes.

The clathrin photoactivation and photoconversion experiments in Fig. 1 were done using a setup that has been described previously in detail (Roy et al., 2012). All other photobleaching and photoconversion experiments were done on the Nikon set-up described above, using Andor Mosaic3 Digital Micromirror Device (DMD) attached to a 405 nm diode laser (450 mW). All synaptic photobleaching and photoconversion experiments were done on DIV-13–14 neurons – a time when synapses are established in our post-natal cultures. For photobleaching, *en passant* boutons were bleached using the 405 nm laser (5 consecutive pulses of 1.5s each), and imaged at every 2s for several minutes. For photoconversion, Dendra2:CLC at boutons was photoconverted by pulsing the 405 nm laser for 1 sec, and boutons were imaged at 2s time-intervals for several minutes. For the single clathrin-packet experiments, boutons were imaged every 1s for several minutes. All movies were analyzed using kymographs generated with a built-in function in MetaMorph (Molecular Devices, LLC). Photoactivation and photoconversion data was analyzed using the intensity-center/bin-center shift method described previously in several publications from our group (Chakrabarty et al., 2019; Ganguly et al., 2017; Roy et al., 2012; Scott et

al., 2011; Tang et al., 2013). All statistical analysis was performed using Graph Pad Prism (Graph Pad Software, San Diego, CA).

Rapamycin-inducible FKBP-FRB dimerizer and FM4-64 assays—Primary hippocampal neurons were co-transfected with the FRB-GFP:CLC and FKBP-BicD2 (+/– Tdtomato-tagged) constructs. Experiments were performed on 9–15 DIV neurons. Live neurons were incubated with 100 nM Rapalog for indicated times. Subsequently, neurons were either used for FM4-64 uptake assay or fixed for immunocytochemistry. Viability of neurons was evaluated using dead cell stain (Live-or-Dye NucFix™ Red), following the manufacturers' protocol. In brief, after the FKBP-FRB experiment, live neurons were incubated with the stain (0.1x in PBS) for 10 mins at 37°C. Thereafter, neurons were washed with PBS and imaged in HELF media at 37°C. For positive controls, neurons were treated with 0.01% Triton for 1 min before staining. For the FM4-64 uptake assay, we used a protocol specifically optimized to evaluate CME (Cousin et al., 2018). Briefly, neurons were washed with imaging buffer (2.5 mM KCl, 136 mM NaCl, 1.3 mM MgCl₂, 2 mM CaCl₂, 10 mM glucose, 10 mM HEPES) followed by 30 s incubation in high potassium stimulation buffer (50 mM KCl, 86 mM NaCl, 1.3 mM MgCl₂, 2 mM CaCl₂, 10 mM glucose, 10 mM HEPES) containing 10 μM FM4-64. Thereafter, neurons were washed with imaging buffer containing 10 μM FM4-64 for 90s. Finally, neurons were washed with Imaging buffer containing 1mM Advasep-7 for 2 min and imaged. All experiments were done at room temperature, with pH of all buffers adjusted to 7.4, and supplemented with 10 μM CNQX, 50 μM DAP-5. For some experiments, neurons were preincubated with 80 μM Dynasore before FM4-64 uptake.

Biochemical fractionation, western blotting and immuno-precipitation from mouse sciatic nerves—In vivo biochemical assays were adapted from protocols described earlier by (Das et al., 2013; Ganguly et al., 2017; Scott et al., 2011; Tang et al., 2012). Briefly, sciatic nerves were dissected from 6–8-wk-old mouse CD1 (WT) mouse. 64 sciatic nerves were pooled for each round of immune-precipitation (IP) and crushed in liquid nitrogen using a motor pestle and then homogenized in nondenaturing buffer (1X IP buffer, Invitrogen Dynabeads™ Co-Immunoprecipitation Kit [ThermoFisher Scientific, USA]) using 18 G and 23 G needles in the presence of protease inhibitor cocktail (Sigma-Aldrich). The resulting homogenate was centrifuged at 1,000 g for 20 min at 4°C to obtain a nuclear pellet (P1) and a post-nuclear supernatant (S1). The S1 supernatant was then centrifuged at 10,200 g for 20 min at 4°C to obtain a crude synaptosomal fraction (P2) and synaptosome-depleted fraction (S2). IP was performed using the DynaBeads Co-Immunoprecipitation kit (14321D; Thermo Fisher Scientific). After centrifugation, the S2 fraction was divided equally and incubated with 10.5 mg anti-Clathrin-Heavy-Chain antibody (ab21679; Abcam, Cambridge MA) and anti-Mouse IgG (ab190475; Abcam, Cambridge MA) for overnight coupling with magnetic beads at 37°C. All following washes were performed as per the manufacturer's protocol. The S2 fraction was incubated with the antibody-coupled beads for 35 minutes at 4°C on a rotor. After the final wash, 3 mg beads from each fraction were subjected to 2D gel electrophoresis on a 4–12% gradient SDS-page gel and then probed with the anti-Clathrin-Heavy-Chain antibody to determine the efficacy of immunoprecipitation. The remaining 7.5 mg of S2 fraction lysate beads were then used

for MudPIT-MS analysis. Two independent repeats were performed with sciatic nerve axons S2 lysate.

Protein identification through MudPIT-MS analysis and in silico data analysis

—Overall steps in MudPIT-MS protocols were identical to our previous studies (Ganguly et al., 2017). Briefly, beads coated with the clathrin heavy chain antibody (or mouse IgG) were dissolved in 100 μ l of 8 M urea in 100 mM Tris, pH 8.5, followed by reduction and alkylation in 10 mM Tris (2-carboxyethyl) phosphine hydrochloride (Sigma) and 55 mM iodoacetamide (Sigma-Aldrich), respectively. This was followed by digestion in trypsin (Promega; incubated at 37°C overnight in the dark) and magnetic bead removal by a magnetic separator. The resulting protein digest was acidified in formic acid followed by centrifugation at 14,000 rpm for 10 min. Thereafter, the supernatant was pressure loaded onto a 250 μ m inner diameter–fused silica capillary column (Polymicro Technologies). This column was fitted with a Kasil frit packed with 2.5 cm of 5 μ m Partisphere strong cation exchange resin (Whatman) and 2.5 cm of 5 μ m C18 resin (Phenomenex). After desalting, this biphasic column was connected to a 100 μ m inner diameter–fused silica capillary (Polymicro Technologies) analytical column with a 3 μ m pulled tip packed with 10 cm of 3 μ m C18 resin (Phenomenex). The entire three-phase column was then laced in line with a 1,200 quaternary HPLC pump (Agilent Technologies) and analyzed using a modified 12-step separation described previously (Washburn et al., 2001). As peptides were eluted from the microcapillary column, they were electrosprayed directly into a hybrid LTQ Orbitrap Velos mass spectrometer (Thermo Fisher Scientific). A cycle consisted of one full-scan mass spectrum (300–1,600 m/z) followed by 20 data-dependent collision-induced dissociation tandem MS spectra. The application of mass spectrometer scan functions and HPLC solvent gradients was controlled by the Xcalibur data system (Thermo Fisher Scientific). Tandem MS spectra were extracted using RawXtract (version 1.9.9; (McDonald et al., 2004) and searched with the ProLuCID algorithm (Xu et al., 2015) against a mouse UniProt database concatenated to a decoy database in which the sequence for each entry in the original database was reversed (Peng et al., 2004). The ProLuCID search was performed using semienzyme specificity and static modification of cysteine because of carboxyamidomethylation (57.02146). ProLuCID search results were assembled and filtered using the DTA Select algorithm (version 2.0; (Tabb et al., 2002)). The protein identification false positive rate was kept below 1%, and all peptide–spectra matches had <10 ppm mass error.

For selection of peptides from the raw MudPIT data, peptides were subjected to the following selection criteria. First, the same peptide fragment had to appear in both rounds of the co-immunoprecipitation, followed by the criteria that the total raw spectrum count from both rounds for the peptide should be >30. Finally, only peptides with combined spectrum counts twenty-fold or higher than the combined spectrum counts from IgG coupled beads were selected. Only peptides that meet all three criteria were included in the final list (Supp. Table 1). The identification of peptides and gene names was performed using the Uniprot database (<https://www.uniprot.org/>). For generation of the clathrin protein interaction map from the S2 fraction all known interactions from experimental data were determined from the String protein interaction database (<http://www.string-db.org>).

Sample preparation, data acquisition and data processing for electron

tomography—For EM, mouse hippocampal neurons were plated at a density of 25,000 cells/100 μ l in wells of gridded MatTek dishes coated with poly-D-lysine. DIV 9–10 neurons were transfected with Apex:GFP:CLC using lipofectamine 2000 as described previously (Ganguly et al., 2017). 6–8 hours before transfection, 7 μ M Hemin chloride (Sigma-Aldrich) was added to the culture medium and kept in the medium until the fixation step (below). 12–16 hours after transfection, transfected neurons were identified based on the GFP signal, and corresponding grid numbers were noted. Following this, neurons were fixed for 5 mins at room temperature, and then 60 minutes on ice in 2.5 % glutaraldehyde in 0.1M sodium cacodylate buffer (pH 7.4). After washes in cacodylate buffer and a quenching step in 20mM glycine, cells were incubated with freshly prepared DAB (25.24 mM) and 0.03% H₂O₂ for 25–30 minutes. Neurons were then washed in 0.1M cacodylate buffer and post-fixed with 1% osmium tetroxide for 30 minutes on ice. The neurons were washed with ddH₂O and then dehydrated and embedded in Durcupan epoxy resin. After curing, the coverslip from the bottom of the MatTek dish was removed gently, and grid numbers with transfected neurons were sawed out and mounted on a dummy acrylic block for sectioning. Ultra-thin (70 nm) and semi-thin sections (200 nm) were cut using a diamond knife (Diatome). To improve stability of specimens under the beam of the EM for tomography, these sections were coated with carbon on both sides. Colloidal gold particles (5 nm and 10 nm diameter) were deposited on each side of the sections to serve as fiducial markers.

Ultra-thin sections were imaged on the JOEL 120 kV while the EM tomogram data were obtained from distal axonal regions (>150 μ m from the cell body) of transfected neurons using a FEI Titan high base microscope operated at 300 kV and micrographs were produced using a 4K \times 4K Gatan CCD camera (US4000). Both microscope and detector were controlled by the Serial EM software package (Mastrorade, 2005) which managed the automated tilt series acquisition. Technical details of how the sections of the tomogram were acquired, image series were aligned properly, and its 3D representations were created have been described by (Phan et al., 2017). Briefly, sections were tilted every 1° from –60° to +60°, aligned properly using colloidal gold particles as fiducial markers and final high-quality 3D representations were built from the projection sets using a custom written non-linear bundle adjustment scheme in TxBR (Lawrence et al., 2006). 3D tomograms were visualized in IMOD (Kremer et al., 1996) and only axon regions with clathrin organelles which do not open to the surface were selected for representative images. EM tomograms were performed on Apex-GFP:CLC transfected neurons from three separate cultures. At least 2–3 independent transfected neurons from each experiment were imaged. 8 axons and 9 dendrites were imaged in total, with tomograms for at least 3 ROIs acquired for each transfected axon and dendrite. A total of 62 coated-pits were identified from tomograms of dendrites, and 17 clathrin coated structures were identified in tomograms from axons. For immuno-EM experiments, a mouse monoclonal antibody against clathrin (clone X22, 1:200–500; Affinity Bioreagents) was used. Detailed procedures including preparation, fixation and immunogold labeling are described in Tao-Cheng, 2020.

DNA-PAINT imaging and image analysis—Rat hippocampal neurons were cultured on 18 mm coverslips at a density of 6,000/cm from embryonic day 18 pups following

established guidelines of the French Animal Care and Use Committee (French Law 2013–118 of 1st February 2013) and approval of the local ethics committee (agreement 2019041114431531-V2 #20242). To selectively label identified axons, neurons in some experiments were transfected the day before fixation with actin-GFP using Lipofectamine 2000 according to the manufacturer's instructions. After 9 to 13 DIV, neurons were fixed at RT in phosphate buffer (PB) 0.1M pH 7.3 containing 4% paraformaldehyde and 4% sucrose for 10 minutes, followed by permeabilization and blocking in incubation buffer (IB): 0.22% gelatin, 0.1% Triton in PB 0.1M pH 7.3 at RT for 2 hours, and incubated overnight at 4°C with primary antibody in IB: polyclonal chicken anti-Map2 (Abcam #53392, 1:1000), polyclonal rabbit anti-clathrin heavy chain (Abcam #21679, 1:150) and monoclonal mouse anti β 2-spectrin (for axon shaft images, BD Bioscience 612563, 1:150) or monoclonal mouse anti-synapsin (Synaptic Systems #106–001, 1:500. After rinses, neurons were incubated with secondary antibodies, including anti-rabbit D2 and anti-mouse D2 DNA-coupled secondary antibodies for DNA-PAINT (Ultivue) at RT for 1 hour.

DNA-PAINT imaging (Jungmann et al., 2014) was performed on an N-STORM microscope (Nikon Instruments) in imaging buffer with 0.25–1 nM Imager-650 and nM Imager-560 (Ultivue). The sample was alternatively illuminated at 647 nm and 561 nm (full laser power) and 20,000–30,000 images of each channel were acquired at 25–33 Hz. DNA-PAINT acquisitions were processed using the N-STORM software followed by post-processing; filtering and image reconstruction at 4 or 12 nm/pixel with the ThunderSTORM Fiji plugin (Jimenez et al., 2019). For 3D rendering, localizations were density-filtered (50 nm 3D radius) before reconstruction of individual presynapses as of 4-nm voxel stacks. Stacks were then 3D gaussian-smoothed and imported into ChimeraX software (Goddard et al., 2018). After surface thresholding and smoothing (default values), 3D rendering images and movies were exported as images and movies. Clathrin particles were segmented using the Segger plugin and their volume measured. Volume V in μm^3 was converted in equivalent sphere diameter D in nm using $[D=1000*2*V^{(1/3)}*3/(4*\pi)]$ for graphing. For presynaptic clathrin particles, only particles contacting or apposed to the synapsin cluster were counted and measured.

QUANTIFICATION AND STATISTICAL ANALYSIS

FRAP Quantification—FRAP data were analyzed as previously described (Applewhite et al., 2007; Boyer et al., 2010). Background fluorescence values were subtracted from all data at each time point, and photobleaching was estimated by fitting an exponential decay function ($F = F_0 \times e^{-kt}$, where F is fluorescence, F_0 is initial fluorescence, k is the decay time constant, and t is time) to the fluorescence of an unbleached control region in the same image. Fluorescence curves were normalized between cells by subtracting the fluorescence immediately after bleaching (time = 0), and then dividing by the average of the 5 pre-bleach fluorescence time points. Fluorescence recovery half-time and immobile fraction were calculated from an inverse exponential decay ($F = A \times (1 - e^{-t/\tau})$, where F is fluorescence, A is the recovery plateau fluorescence, τ is the recovery time constant, and t is time) fit to the average fluorescence recovery curve for each protein or treatment. Photobleaching curves were fit to the data using the Solver add-in of Microsoft Excel 2013, and fluorescence recovery curves were fit using GraphPad Prism.

Statistical Analyses—Datasets were tested for normality using the Shapiro-Wilk W test with α set at 0.05. Pairwise comparisons were made using a Student's *t*-test only when both groups being compared were found to be normally distributed: in all other cases, a Mann-Whitney nonparametric U test was used. Correlation of clathrin to synapsin volume at synapses was performed using a Pearson's test. All three-group comparisons contained non-normally distributed data; therefore, nonparametric Kruskal-Wallis tests were used for all three-group comparisons. Post-hoc pairwise comparisons were performed with Mann-Whitney nonparametric U tests and were corrected for multiple comparisons using the Benjamini-Krieger-Yekutieli two-stage step-up method. No pre-hoc sample size estimates were determined prior to experiments. Details of experimental design are described in the experiment specific subsections of Method details section. All the data are presented as mean \pm SEM. The values of *n* and what *n* represents are included in the legends of the respective figures.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Highlights:

- Clathrin moves as stable ‘transport-packets’ in axons with overall anterograde bias
- Stable clathrin-packets switch to a dynamic state upon reaching synapses
- Synaptic clathrin-packets are radially organized around synaptic vesicle clusters
- Depletion of synaptic clathrin-packets impairs synaptic vesicle recycling

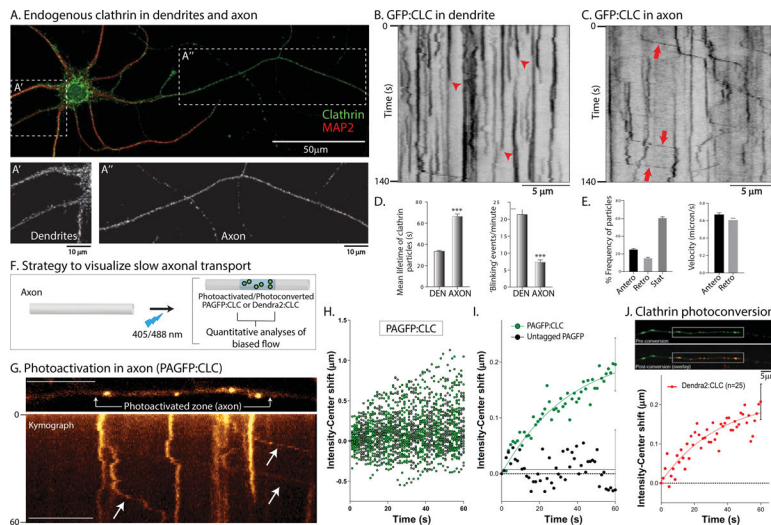


Figure 1: Differential dynamics of clathrin in dendrites and axons.

(A) Clathrin and MAP2 immunostaining in cultured hippocampal neurons. Note punctate clathrin in both somatodendritic (A') and axonal (A'') compartments.

(B) Kymograph of GFP:CLC in a dendrite. Note abrupt appearance and disappearance of fluorescence (red arrowheads), indicating assembly/disassembly of clathrin in coated-pits (CME). Scale bars=5 μ m.

(C) Kymograph of GFP:CLC in an axon. Note rapid, infrequent, and anterogradely-biased movement of clathrin particles (some marked by red arrows; more examples in Supp. Fig. 1A). Scale bars=5 μ m.

(D) Mean lifetimes of GFP:CLC particles – as determined from kymographs – were significantly higher in axons, while ‘blinking events’ – representing clathrin assembly/disassembly and receptor mediated endocytosis is much lower in axons, compared to dendrites (1113 particles from 22 dendrites and 403 particles from 41 axons were analyzed; data was pooled from 3 independent experiments; *** $p < 0.0001$)

(E) About 40% of the axonal GFP:CLC particles were mobile, with a larger fraction of particles moving anterogradely (460 anterograde and 263 retrograde events were analyzed from 41 axons; data was combined from 3 independent experiments).

(F) Strategy for analyzing biased axonal transport of clathrin in axons (see results for details).

(G) Kymograph from a photoactivation (PAGFP:CLC) experiment. Note anterograde movement of photoactivated clathrin particles (arrows). Scale bars=10 μ m.

(H) Raw data of intensity-center shifts from all PAGFP:CLC experiments in axons. Note anterograde drift of datapoints (38 neurons from 3 separate cultures were analyzed).

(I) Ensemble data showing mean intensity-center shift of PAGFP:CLC and PAGFP-only in axons, with fitted curves. While there is an overall anterograde bias of PAGFP:CLC, there is no bias with untagged PAGFP-only (10 neurons from 2 separate cultures were analyzed for PAGFP-only).

(J) Neurons were transfected with Dendra2:CLC, axonal clathrin particles were identified by green fluorescence, and intensity-center shifts of photoconverted (red) particles was

analyzed. Note anterograde shift of the particles (25 neurons from 3 separate cultures were analyzed). Scale bars=5 μ m.

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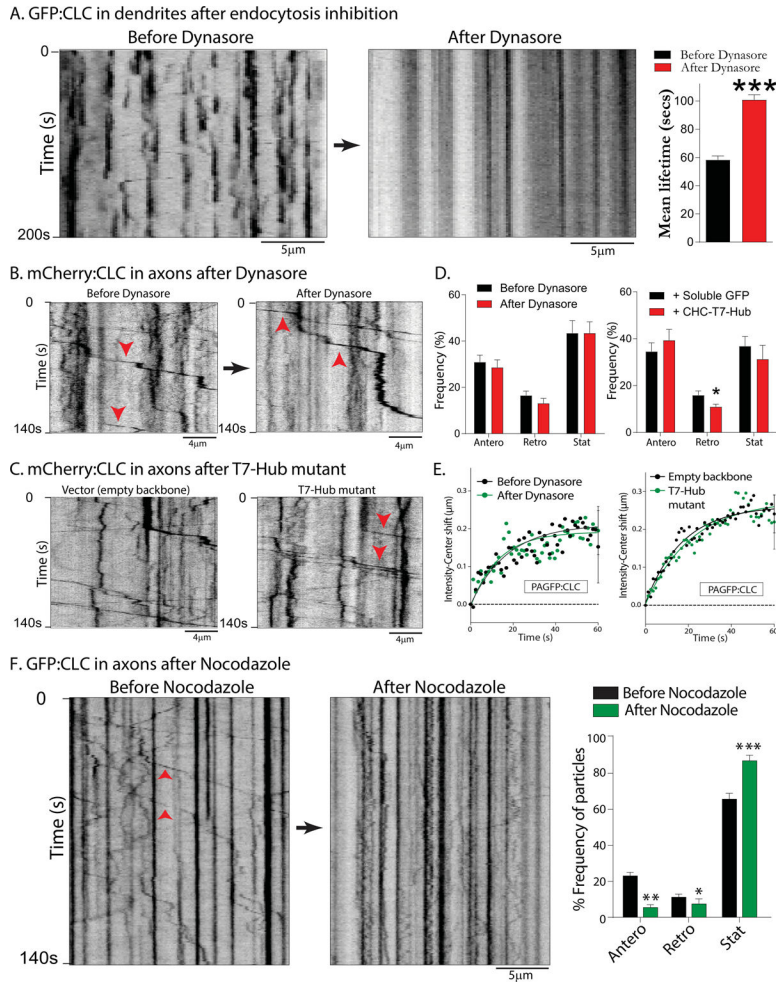


Figure 2: Motility of axonal clathrin transport-packets is independent of endocytosis, but dependent on microtubules.

(A) Kymographs of GFP:CLC from a dendrite before and after endocytosis inhibition (Dynasore). Note that blinking events – reflecting endocytosis – are abolished after drug-treatment. GFP:CLC fluorescence-lifetimes are quantified on right (~ 150–250 particles from 10 neurons were analyzed; data from three separate cultures; $p < 0.0001$) Scale bars=5 μm .

(B) Kymographs of mCherry:CLC from same axon before and after Dynasore. Note that vectorial movement of clathrin transport-packets (some marked by arrowheads) continue after drug-treatment. Scale bars=4 μm .

(C) Kymographs of axonal mCherry:CLC from neurons transfected with a dominant-negative clathrin mutant (T7-Hub) that specifically interferes with clathrin-dependent endocytosis (see Supp. Fig. 2); or control. Note that vectorial movement of clathrin transport-packets (some marked by arrowheads) are similar in the two groups. Scale bars=4 μm .

(D) Quantification of all mCherry:CLC axonal transport data – with and without endocytosis inhibition – global or clathrin-dependent. (~ 140–170 motile particles were analyzed from 10–12 neurons; data from 2–3 separate cultures; * $p < 0.01$)

(E) Ensemble intensity-center shift data (PAGFP:CLC slow transport assay) from neurons treated with Dynasore or co-transfected with the T7-Hub mutant. Note that inhibiting global – or clathrin-dependent – endocytosis has no effect on the slow anterograde bias of the clathrin population in axons (20–30 axons from two and four separate cultures were analyzed).

(F) Kymographs of GFP:CLC dynamics from the same axon before and after treatment with 10 μ g/ml Nocodazole for 30 minutes; quantification on right. Note that motility of the clathrin transport-packets is attenuated after drug-treatment. Scale bars=5 μ m.

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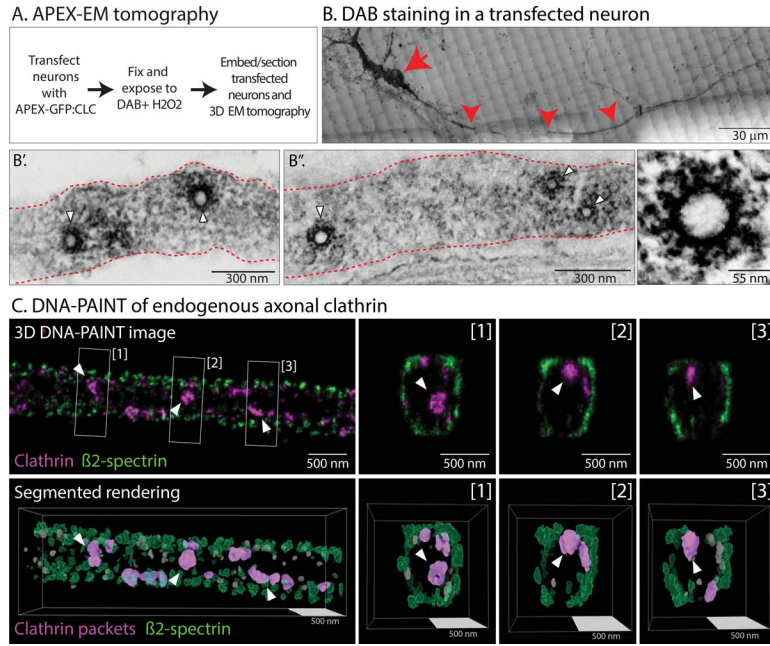


Figure 3: Ultrastructure and super-resolution-imaging of axonal clathrin transport-packets. (A) Strategy to visualize clathrin by Apex-labeling and EM-tomography. (B) A DAB-stained neuron identified on gridded membrane (soma and axon marked by arrow and arrowhead respectively; also see methods and Supp. Fig. 3). Scale bars=30 μm. (B' and B'') EM of the axon above (a single plane of tomogram shown). Note clathrin-coated structures in the axon-shaft (white arrowheads), with average diameters of ~ 50 nm (spherical core). Representative 3D-EM from three separate cultures (see “Methods” for more details). Scale bar (lower left) for B' is 300 nm, B'' is 300 nm and lower right is 55 nm. (C) DNA-PAINT of endogenous clathrin in the axon shaft. **Top panels:** β2-spectrin (green, to mark axonal boundaries) and clathrin heavy chain (magenta). An X-Y image is shown on the left, along with three Z-slices on right ([1]-[3]). **Bottom panels:** Segmented rendering from 3D-PAINT images (small excluded particles are in grey). Equivalent sphere diameters for the clathrin particles marked by arrowheads: 79, 70 and 70 nm for particles highlighted in [1], [2] and [3] respectively. Scale bars=500 nm.

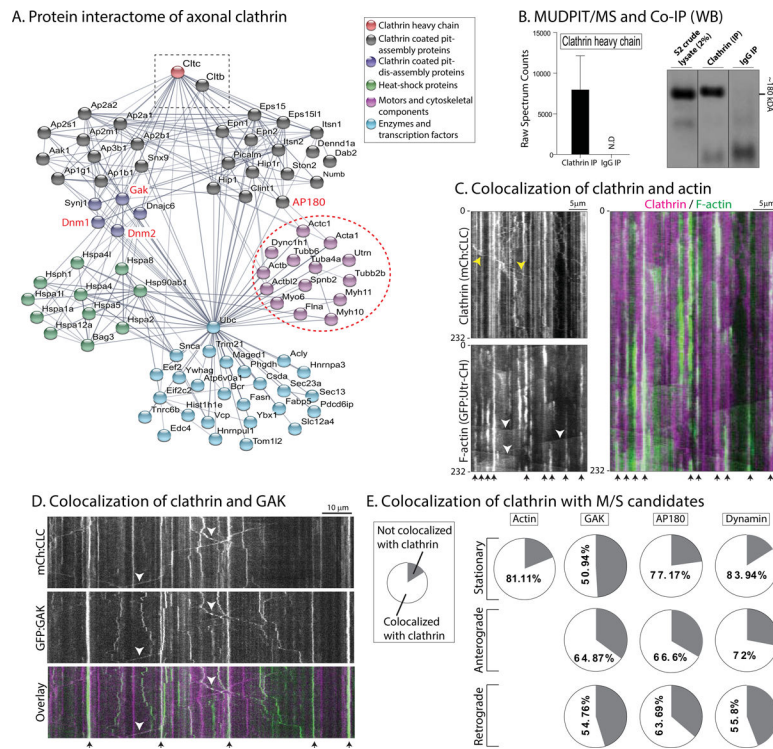


Figure 4: The axonal clathrin interactome.

(A) An interactome of clathrin-associated proteins (clathrin marked by dashed rectangle), grouped into functional clusters. The ‘actin-hub’ (dashed circle) was of particular interest as previous studies have established mechanistic links between clathrin and actin (also see results).

(B) High clathrin spectrum-counts in MS-data, confirmed by western blots showing clathrin in IP fractions.

(C) Kymographs from axons of neurons co-transfected with mCherry:CLC and a GFP-tagged probe for F-actin (GFP:Utr-CH). Note extensive colocalization of stationary clathrin and actin (arrowheads at the bottom, merged kymograph on right). Moving clathrin particles and “actin trails” are marked on the kymographs by yellow and white arrowheads respectively. Time in seconds is marked on the left of the kymographs, and scale bar on upper right. Scale bars=5 μ m.

(D) Kymographs from axons of neurons co-transfected with mCherry:CLC and GFP:GAK. Note colocalization of stationary particles (arrowheads at the bottom of kymographs), and co-transported clathrin and GAK particles (arrowheads). Time in seconds is marked on the left of the kymographs, and scale bar on upper right. Scale bars=10 μ m.

(E) Quantification of kinetic data from simultaneous imaging of clathrin with candidates from M/S data (GAK, AP180 and dynamin). Note that a substantial amount of clathrin particles were co-transported with GAK, AP180 and dynamin (~ 140–340 particles were analyzed from 12–19 neurons; data from 2–3 separate cultures).

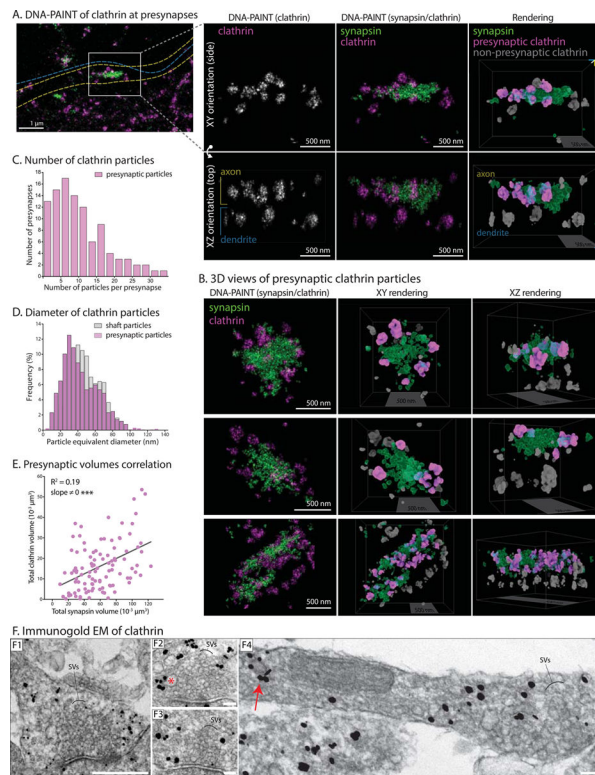


Figure 5: The nanostructure of clathrin at the presynapses.

(A) **Left:** DNA-PAINT of endogenous clathrin (magenta) and synapsin (green, SV-marker) at an *en-passant* bouton. Outline of a dendrite (blue dashed line) and axon (yellow dashed line) are shown (see full image in Supp. Fig. 5A). Scale bars=1 μ m.

Right: 3-D views of rectangular inset with XY and XZ (rotated) views; along with 3-D rendering and segmentation (extreme right panels). The rendered view shows presynaptic clathrin in magenta, non-presynaptic clathrin in grey, and synapsin in green. Note clathrin particles abutting the presynaptic SV-cluster. Also see Supp. Movie 4. Scale bars=500 nm. (B) 3D views of more presynapses showing DNA-PAINT image and segmented rendering (also see Supp. Movie 4). Scale bars=500 nm.

(C) Histogram of segmented presynaptic clathrin particles per presynapse (1261 clathrin particles from 102 synapses analyzed).

(D) Overlaid histograms showing frequency distribution of presynaptic clathrin particle-diameters (1261 particles from 3 independent experiments), compared to clathrin particle-diameters in axon shaft (685 particles from 3 independent experiments).

(E) Total clathrin volume (cumulative of all clathrin particles) at a given bouton correlated with the total volume of synapsin.

(F) Immuno-gold EM of clathrin (F1-F4; F1-F4 – silver enhancement of nanogold particles). Note most particles are located at the periphery of SV-clusters, and some are associated with cisternae that may be endosomes (red asterisk in F2 for example). F4 shows segment of an axon with presynapse, and red arrow marks a coated structure resembling axonal transport-packet. Note that the labeling is specific for clathrin; also see Supp. Fig. 5B Scale bar (lower right) for F1 is 500 nm and for F2-F4 is 100 nm.

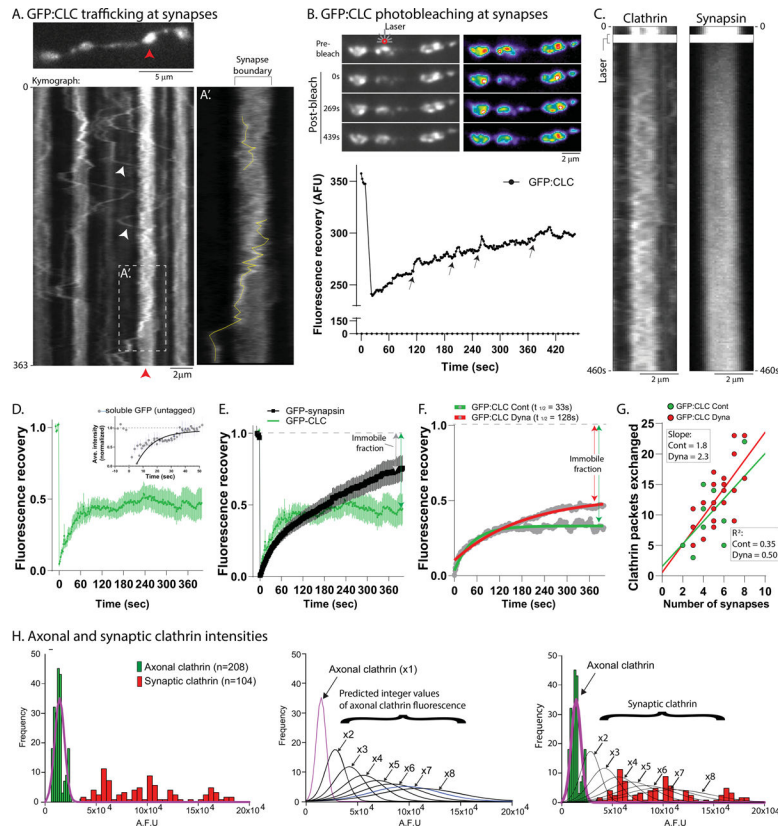


Figure 6: Transported clathrin-packets are captured and immobilized at presynapses.

(A) Rapid imaging of GFP:CLC around boutons, first frame on top and kymograph at bottom. Note that clathrin particles are rapidly transported between boutons and often localize to synapses (white arrowheads, targeting to synapse marked by red arrowhead). (A') is a zoom of the boxed region in kymograph showing trajectories of two particles. Scale bar for upper left is 5 μm and lower left is 2 μm .

(B) GFP:CLC was bleached in a single bouton and fluorescence recovery was monitored over time, as shown in the time-series greyscale and pseudocolor-heatmap images. Note gradual recovery of fluorescence over time, also evident in the FRAP curve below (small arrowheads point to transient peaks due to particle-entry into bleached bouton; also see Supp. Movie 5). Scale bars=2 μm .

(C) Bouton kymographs from FRAP of GFP:CLC and GFP:synapsin. Note that while fluorescence recovery of clathrin is particulate, the rise of synapsin fluorescence is diffusive. Scale bars=2 μm .

(D) Bouton FRAP recovery of GFP:CLC was incomplete, suggesting an immobile pool of clathrin (FRAP of soluble, untagged GFP in boutons recovered completely, inset). For GFP:CLC, 12 neurons were analyzed from 3 separate cultures.

(E) Comparison of GFP:CLC and GFP:synapsin bouton FRAP. Note that clathrin has a larger immobile pool than synapsin (12–14 neurons from 2–3 separate cultures were analyzed).

(F) Blocking endocytosis with Dynasore attenuated the immobile fraction of GFP:CLC in bouton-FRAP experiments (i.e. increased mobility of clathrin-packets), indicating that

FRAP-recovery was not due to clathrin on newly-generated endosomes at adjacent synapses (11–15 neurons from 3 separate cultures, for each condition). For each experiment, same coverslip was evaluated before/after Dynasore to minimize variability.

(G) More GFP:CLC particles exchanged between synapses after Dynasore-treatment (moved from one synapse into another and stayed there for at least a minute). ~ 200–400 particles moving between ~ 100 synapses (16–21 neurons per condition) were analyzed.

(H) Integers of axonal clathrin-packets localize to synapses. **Left:** Histogram showing distribution of GFP:CLC intensities in axons and synapses. Note that while axonal clathrin intensities are tightly clustered in a single peak (green), synaptic intensities are distributed as multiple peaks of fluorescence (red). **Middle:** Mean fluorescence of axonal clathrin was considered as 1x, and hypothetical integer-multiple curves were generated (fitted curves shown). **Right:** Axonal and synaptic GFP:CLC fluorescence data were overlaid with the curves shown in middle panel. Note that data predicts that the synaptic clathrin fluorescence is composed of 4–8+ integer-multiples of axonal fluorescence (100–200 particles were analyzed from three separate cultures).

Key for kymographs: elapsed time in seconds on left or right and scale-bar on lower right (laser illumination also marked).

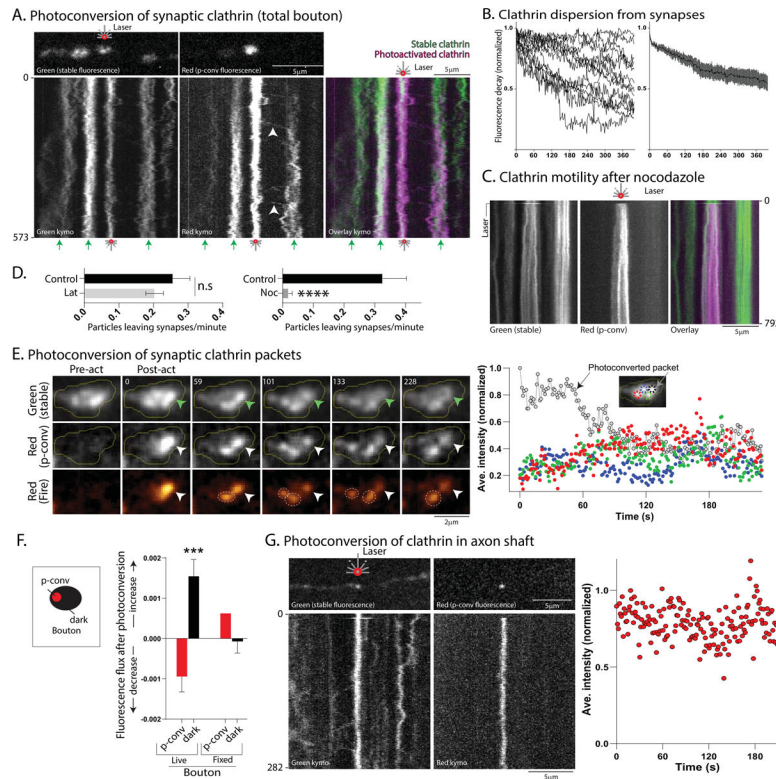


Figure 7: Dynamics of synaptic clathrin-packets.

(A) First post-photoconverted frames (above) and kymographs (below) from photoconversion of Dendra2:CLC at a single bouton. Photoconverted bouton is marked by red laser symbol, and non-photoconverted boutons by small green arrows at bottom. Note that over time, fluorescent clathrin particles leave the single photoactivated bouton and are immobilized at adjacent boutons (white arrowheads in middle kymograph).

(B) Decay of photoconverted Dendra2:CLC red fluorescence from single boutons (raw and ensemble data; 13 neurons from 2 separate cultures).

(C) Nocodazole (10 μ g/ml for 40 minutes) attenuated export of synaptic clathrin particles in single-bouton photoconversion experiments while Latrunculin (100nM for 15 minutes) had no significant effect; quantified in D (15–20 neurons from 2–3 separate cultures analyzed for each control- and drug-treated group).

(E) **Left:** Dendra2:CLC was photoconverted in a single synaptic clathrin-packet (arrowhead, elapsed time in seconds on top left). Note that after photoconversion (middle and bottom panels), the photoconverted packet lost fluorescence, while there was an increase of fluorescence in adjacent packets (identified by stable green fluorescence, some marked by dashed circles in bottom panel). **Right:** Quantification of intensities in individual packets (normalized to highest post-conversion intensity). Colors in graph correspond to the colors of circles marking packets in inset-image. Black dashed circle marks the green clathrin packet that was photoconverted to red. Note reciprocal fluctuations in photoconverted and non-photoconverted packets. Scale bars=2 μ m.

(F) Quantification of fluorescence change after single-packet photoconversion in boutons. Schematic on left shows general principle of quantifying photoconverted and non-photoconverted (dark) regions; see “methods” for details. Note loss of intensity from

photoconverted packet within bouton (red bar, left), with a corresponding gain in adjacent non-converted regions (black bar, left). Similar reciprocal changes in fluorescence were not seen after fixing the neurons, though the photoactivation still occurred as expected (red/black middle bars; 6 neurons from 1 culture for fixed boutons and 10 cells from 2 separate cultures for live boutons).

(G) Photoconversion of single Dendra2:CLC clathrin-packet in axon. Note no change in fluorescence, quantified on right (8 neurons from 2 separate cultures analyzed). Scale bars=5 μm .

Time in seconds on left or right of kymograph and scale bar on upper right. **** $p < 0.0001$, *** $p < 0.001$. See Supp. Fig. 7 for more examples.

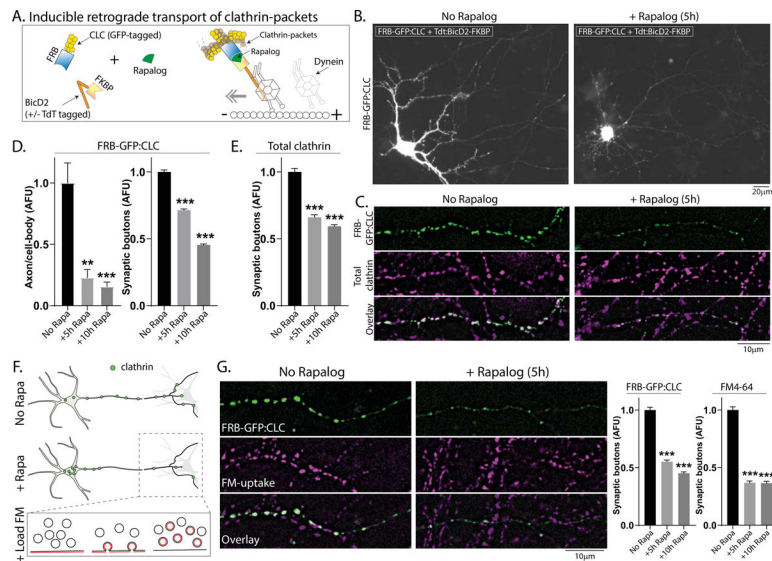


Figure 8: Depletion of synaptic clathrin packets impairs function.

(A) Principle: GFP:CLC was tagged to FRB, and the dynein adaptor BicD2 was tagged to FKBP (+/- TdTomato). Upon adding Rapalog, FKBP and FRB – along with clathrin and BicD2 – would dimerize, transporting clathrin-packets into the soma and depleting them from synapses.

(B) Representative images showing clustering of FRB-GFP:CLC in cell-bodies after Rapalog treatment (constructs used on top left of images). Also note that transfected FRB-GFP:CLC extensively colocalizes with endogenous clathrin (Supp. Fig. 8). Scale bars=20 μ m.

(C) Representative images showing that upon Rapalog-treatment, both tagged and endogenous clathrin is depleted from boutons – quantified in (D-E; ~ 900–1800 boutons were analyzed for each condition from 3 separate coverslips) Scale bars=10 μ m.

(F) Schematic showing plan of FM-dye uptake experiments (to evaluate SV-recycling) in clathrin-depleted boutons.

(G) Representative images (left) and quantification (right) showing the reduction of both clathrin and FM-uptake in Rapalog-treated neurons at room temperature (~ 600–950 boutons were analyzed for each condition from 3 separate cultures) – similar results were seen at 37°C, see Supp. Fig. 8E. ** $p < 0.01$, *** $p < 0.001$. Scale bars=10 μ m.

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Anti-Clathrin heavy chain antibody	Abcam	Cat#ab21679; RRID:AB_2083165
Chicken polyclonal anti-GFP antibody	Abcam	Cat#ab13970; RRID:AB_300798
Anti-rabbit Alexa Fluor 594	Invitrogen	Cat#A11037; RRID:AB_2534095
Anti-Chicken Alexa Fluor 488	Invitrogen	Cat#A11039; RRID:AB_142924
polyclonal chicken anti-Map2	Abcam	Cat# ab5392; RRID:AB_2138153
monoclonal mouse anti β 2-spectrin	BD Bioscience	Cat#612563; RRID:AB_399854
Anti-Clathrin mouse monoclonal antibody	Affinity Bioreagents	Cat#MA1-065; RRID:AB_2083179
Anti-Mouse IgG	Abcam	Cat#ab190475; RRID:AB_2827162
monoclonal mouse anti-synapsin	Synaptic Systems	Cat#106-001; RRID:AB_887805
Anti-rabbit D2 (Ulvivue Kit)	Ulvivue	Cat#Ulvivue-2
Anti-mouse D2 (Ulvivue Kit)	Ulvivue	Cat#Ulvivue-2
Chemicals, peptides, and recombinant proteins		
Dynasore	Abcam	Cat#ab120192
Latrunculin A	Life Technologies	Cat#L12370
Nocodazole	Millipore-Sigma	Cat#487928-10MG
Swinholide A	Millipore-Sigma	Cat#574776-10UG
A/C heterodimerizer	Takara	Cat#635056
FM4-64	Invitrogen	Cat#T13320
Advasep-7	Millipore-Sigma	Cat#A3723
HBSS	GIBCO	Cat#14025092
6-cyano-7-nitroquinoxaline-2,3-dione CNQX	Tocris Biosciences	Cat#1045
D,L-2-amino-5-phosphonovaleric acid AP5	Tocris Biosciences	Cat#106
VER155008	Tocris Biosciences	Cat#3803
Lipofectamine 2000	Life Technologies	Cat#11668-019
ProFection® Mammalian Transfection System	Promega	Cat#E1200
B27 supplement	GIBCO	Cat#17504044
Glutamax	GIBCO	Cat#35050061
Neurobasal A	GIBCO	Cat#10888022
poly-D-lysine	Sigma	Cat#P0899
Hibernate-E low fluorescence medium	Brainbits	Cat#HE-1f
0.25 % Trypsin-EDTA	Invitrogen	Cat#25200072
Fetal bovine serum	Hyclone	Cat#SH30071.03
Hemin chloride	Millipore-Sigma	Cat#3741
Sodium cacodylate trihydrate	Millipore-Sigma	Cat#C4945
Diaminobenzidine (DAB)	Millipore-Sigma	Cat#D8001

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Osmium Tetraoxide (4% aqueous solution)	Electron Microscopy Sciences	Cat#19150
Paraformaldehyde (16% aqueous solution)	Electron Microscopy Sciences	Cat#15710
Glutaraldehyde (25% aqueous solution)	Electron Microscopy Sciences	Cat#16220
Durcupan ACM, Epoxy Resin (Kit)	Electron Microscopy Sciences	Cat#14040
Tris (2-carboxyethyl) phosphine hydrochloride	Sigma-Aldrich	Cat# 75259
Iodo-acetamide	Sigma-Aldrich	Cat#144-48-9
Protease inhibitor cocktail	Sigma-Aldrich	Cat#P8340
Dynabeads™ Co-Immunoprecipitation Kit	ThermoFisher Scientific	Cat#14321D
1.4 nm Nanogold® secondary antibody	Nanoprobes	Cat#2001
HQ silver enhancement kit	Nanoprobes	Cat#2012-45ML
Critical commercial assays		
Live-or-Dye NucFix™ Red	Biotium	Cat#32010-T
Deposited data		
Proteomics Data	This Paper; ProteomeXchange Consortium (PRIDE database)	Supp.Table 1; PXD027857 (https://www.ebi.ac.uk/pride/)
Experimental models: Organisms/strains		
Mouse: CD1	Charles River Laboratories	Cat#022-CD1
Recombinant DNA		
GFP:CLC	Gaidarov et al.,1999	N/A
pUC57-APEX	Martell et al., 2012	Addgene Plasmid #40306
Dendra2-Lifeact-7	Dendra2-Lifeact-7 was a gift from Michael Davidson	Addgene plasmid #54694
Dendra2:CLC	This paper	N/A
PAGFP-C1	Patterson et al., 2002	Addgene plasmid #11910
PAGFP:CLC	This paper	N/A
CHC-T7-Hub	Liu et al., 1998	N/A
T7-empty backbone	Liu et al., 1998	N/A
GFP:Utr-CH	Burke et al., 2007	Addgene Plasmid #26737
synaptophysin:mCh	Gerrow et al., 2006	N/A
pBa-Flag-BicD2-FKBP	Bentley et al., 2015	Addgene Plasmid #64206
pBa- Flag-tdTM-BicD2-FKBP	Bentley et al., 2015	Addgene Plasmid #64205
pBa-FRB-3myc-Rab5a	Bentley et al., 2015	Addgene Plasmid #64209
FRB-GFP:CLC	This paper	N/A
GFP:synapsin-1a	Gitler et al., 2004	N/A
EGFP-AP180	Bushlin et al., 2008	N/A
GAK-GFP	Massol et al., 2006	N/A
pEGFP-C1-Dynamin	Song et al., 2004	Addgene Plasmid #34680
CLC:mCherry	This paper	N/A
pEGFP-N1	Clontech	Cat#6085-1
Software and algorithms		

REAGENT or RESOURCE	SOURCE	IDENTIFIER
GraphPad Prism	GraphPad Software, La Jolla, CA, USA	Version 7.00; www.graphpad.com ; RRID:SCR_002798
Fiji ImageJ	Schindelin et al., 2012	https://imagej.net/Fiji ; RRID:SCR_002285
MetaMorph Version 7.10	Molecular Devices	https://www.moleculardevices.com/products/cellular-imaging-systems/acquisition-and-analysis-software/metamorph-microscopy ; RRID:SCR_002368
Adobe Illustrator	Adobe	https://www.adobe.com/products/illustrator/free-trial-download.html ; RRID:SCR_010279
RawXtract version 1.9.9	McDonald et al., 2004	http://fields.scripps.edu/downloads.php
N-STORM software	Nikon; Jungmann et al., 2014	https://www.microscope.healthcare.nikon.com/products/super-resolution-microscopes/n-storm-super-resolution ; RRID:SCR_018302
ChimeraX software	Goddard et al., 2018	https://www.rbvi.ucsf.edu/chimerax/ ; RRID:SCR_015872
ThunderSTORM	Jimenez et al., 2019	https://github.com/zitmen/thunderstorm ; RRID:SCR_016897