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In-vivo fast nonlinear microscopy reveals impairment of fast axonal transport induced by molecular motor imbalances in the brain of zebrafish larvae

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Abstract

Cargo transport by molecular motors along microtubules is essential for the function of eucaryotic cells, in particular neurons in which axonal transport defects constitute the early pathological features of neurodegenerative diseases. Mainly studied in motor and sensory neurons, axonal transport is still difficult to characterize in neurons of the brain in absence of appropriate *in vivo* tools. Here, we measured fast axonal transport by tracing the second harmonic generation (SHG) signal of potassium titanyl phosphate (KTP) nanocrystals (nanoKTP) endocytosed by brain neurons of zebrafish (Zf) larvae. Thanks to the optical translucency of Zf larvae and to the perfect photostability of nanoKTP SHG, we achieved a high scanning speed of 20 frames (of $\approx 90 \mu\text{m} \times 60 \mu\text{m}$ size) per second in Zf brain. We focused our study on endolysosomal vesicle transport in axons of known polarization, separately analyzing kinesin and dynein motor-driven displacements. To validate our assay, we used either loss-of-function mutations of dynein or kinesin 1 or the dynein inhibitor dynapyrazole, and quantified several transport parameters. We successfully demonstrated that dynapyrazole reduces nanoKTP mobile fraction and retrograde run length consistently, while the retrograde run length increased in kinesin 1 mutants. Taking advantage of nanoKTP SHG directional emission, we also quantified fluctuations of vesicle orientation. Thus, by combining endocytosis of nanocrystals having a nonlinear response, fast two-photon microscopy, and high-throughput analysis, we are able to finely monitor fast axonal transport *in vivo* in the brain of a vertebrate, and reveal subtle axonal transport alterations. The high spatiotemporal resolution achieved in our model may be relevant to precisely investigate axonal transport impairment associated to disease models.

Keywords: nonlinear nanocrystal, two-photon microscopy, second harmonic generation, single nanoparticle tracking, axonal transport, molecular motor, zebrafish larva, disease models.

Active intracellular transport of cargoes such as organelles, vesicles, RNA granules and other materials, along the cytoskeleton, is an essential molecular process participating in cell homeostasis. This transport is critical in axons and dendrites of neurons, that form extensive branches connecting different organs of the nervous system. Indeed, axonal transport deficits are hallmarks of neurodegenerative diseases¹ and recent studies have also consolidated the key role played by molecular transport in the maintenance and functions of synapses.²

Axonal transport is categorized as fast or slow at velocities ranging from a few $\mu\text{m/s}$ for organelles to $< 0.1 \mu\text{m/s}$ for cytoskeletal proteins,³ respectively. This process takes place along the microtubule (MT) cytoskeleton and is sustained by two types of molecular motors: the superfamily of kinesin motors⁴ ensuring transport of cargoes from MT minus end to the plus end, and the dynein complex⁵ traveling in the opposite direction. As in axons, MT are uniformly polarized with all their plus ends distal from neuron soma, kinesin is responsible for anterograde motions (from soma to axon periphery), and dynein for retrograde displacements (from axon periphery to soma). Over the past few decades, much progress has been made in developing tools and methods to visualize and measure axonal transport.⁶ The most common measurement approach relies on videomicroscopy recording of fluorescently-labeled cargoes displacements. Depending on the type of cargo and the complexity of the biological system (neuronal cell culture, primary neurons *ex vivo* tissue cultures or *in vivo* models), various labeling methods have been selected. The latter include use of live cell compatible dyes like MitoTracker or LysoTracker for mitochondria or lysosomes respectively, fluorescently labeled neurotrophic toxins,⁷ or antibody against neurotrophin⁸ or fluorescent viral particles.^{6,9} Furthermore, taking advantage of genome editing tools applied to model organisms like *Drosophila* and *Danio rerio* (Zebrafish, Zf), fluorescent proteins (FP) have been expressed in specific organelles of given neuronal populations to investigate their axonal transport,^{6,10,11} in different contexts including neurodegenerative diseases,^{12,13} axonal regeneration¹⁴ and age-dependent decline.¹⁵ These physiological models, particularly relevant for the study of complex biological processes by imaging, can be delicate to implement. First of all they rely on transgenesis by injection of the transgene constructs at the single cell level to obtain the mosaic expression¹¹⁻¹³ of the marker of interest, necessary to allow single organelle identification. Then, injected larvae must be selected based on the expression pattern of the marker in terms of localization and level, to be used for axonal transport investigation.

Finally, more challenging studies were also conducted *in vivo* in transgenic mice to monitor the axonal transport of fluorescent mitochondria in peripheral nerves,^{16,17} but they required a complex and invasive surgical exposure of the nerves, animal immobilization and optimization of fast and 3D deep imaging processes.¹⁰ Axonal transport was also measured in neurons of mouse central nervous system (CNS) using two-photon-excited microscopy. This includes neurons of the spinal cord,¹⁸ retinal ganglion cells¹⁹ and cortical neurons.²⁰ However, these approaches not only still require some surgery but photobleaching of the FP reporter and phototoxicity for the animal also limit the temporal resolutions to a maximum of one frame per second, with a spatial resolution of about 200 nm. This time resolution is sufficient to evaluate parameters such as the fraction of mobile cargoes, their velocity and direction of transport but it prevents the observation of short pauses (duration $\lesssim 250$ ms) which are of utmost interest to investigate the microtubule (MT) network complexity and stability,³ that were recently reported by Chowdary *et al.*²¹ and by us in endolysosomal transport in cultured neurons, using a nanoparticle tracing methodology²² acquiring at 20 frame/s (fps).

These micro-pauses were attributed to tug-of-war situations between kinesin and dynein, both attached to the same cargo and pulling in opposite direction, leading occasionally to short pauses when the cargo experiences null net force.²³ The present study aims at accurately investigating axonal transport modifications of endolysosome compartments induced by modulations of molecular motors concentrations in zebrafish larvae brain, *in vivo*. To be sensitive to short pauses, it is therefore critical to achieve a frame rate at least equal to the 20 fps used in sensitive *in vitro* studies.^{21,22} Indeed, intravital imaging of organelle transport can also benefit from the use optically-active photostable nanocrystals internalized in neuronal cells by endocytosis and subsequently traced at high frame rate. This technology was pioneered by Cui *et al*²⁴ who used quantum dot as fluorescent tracers. Later, we reported the use of fluorescent diamond nanocrystals as tracers to reveal subtle changes in endolysosomal transport within cultured neurons established from a transgenic mouse bearing a genetic risk factor of a neuropsychiatric disease.²²

Here we extended such nanoparticle-based assay to measure the endolysosomal transport parameters *in vivo*, in neurons of the brain of intact, living zebrafish larvae. To this aim we used potassium titanyl phosphate (KTiOPO₄, KTP) nanocrystals (size \approx 120 nm) that exhibit large second-order nonlinear optical response, and micro-inject them in optic tectum of Zf larvae. Incidentally, the ability to image these efficient nonlinear KTP nanocrystals (nanoKTP) in live Zf larvae blood circulation at ultra-high frame rate has been recently reported in a wide-field configuration with light-sheet illumination.²⁵ Furthermore, in a previous work, we had shown that nanoKTP are spontaneously internalized in 2D primary cultures of neurons and can be traced by acquisition of their second harmonic generation (SHG) emission under infrared (IR) pulsed laser excitation.²⁶ Key advantages of SHG over fluorescence are non-saturation, non-bleaching, and a reduced background, hence a large signal to background ratio. In addition, the wavelength of SHG emission is tunable and can be adjusted to avoid any overlap with other fluorescent labels used in the same sample.

In the present work, we harnessed these properties combined with fast raster scanning of the IR laser beam to achieve the frame rate of 20 frames/s (fps) for a \approx 80 μ m-sized field-of-view, identical to the one employed in our *in vitro* intraneuronal transport assay.²² Moreover, due to a single dominant coefficient of the nonlinear susceptibility tensor,²⁷ SHG from KTP has a directional emission. We took advantage of this property to investigate the vesicle rotational dynamics during their axonal transport, in a similar way as it was reported in cultured neurons,^{28,29} where very fast dynamics was captured at 500 fps (2 ms frame duration), justifying further our settings to the highest possible frame rate. The amplitude of vesicle rotational motions is an indirect readout of how strongly kinesin and dynein motors tether it to microtubules. The more motors there are, tethering a vesicle to a microtubule, the more constrained are its position and its orientation relative to a reference frame, as the motor counteract Brownian motion. If some motors linked to the vesicle untether from the microtubule, the amplitude of vesicular Brownian motions, that include rotations, will increase. We hypothesize, like in Kaplan *et al.*,²⁹ that the amplitude of orientational fluctuations is a qualitative readout of the number of motors attached to both the endosome and the microtubule, and therefore that it adds significant value to the analysis, at no additional experimental cost.

In our experiment, we showed that the nanoKTP move within axons of periventricular neurons (PVN) that all project radially inside the neuropil to connect retinal ganglion

cells.³⁰ Considering this anatomical characteristics of PVN, we were able to distinguish the anterograde transport, from the retrograde transport, and to quantify associated parameters separately for each direction. While in our previous study in cultured neurons, we had investigated the impact on intraneuronal transport of microtubule disruption or of the level of a kinase phosphorylating MT-associated proteins, without distinction of axons and dendrites, here we focused on active molecular motor-dependent mechanisms. As we addressed specifically axonal transport along polarized MT, we could relate the anterograde or retrograde transport direction to the activity of one type of motors. We considered transgenic Zf engineered to bear loss-of-function alleles of retrograde motor protein Dync1h1 heavy chain 1 of the dynein motor complex³¹ or of the anterograde motor protein Kif5aa.³² As in *dync1h1* mutant all transport was halted, we opted for a drug inhibition strategy to achieve a finer control of retrograde transport, using dynapyrazole, a cell permeable, conformationally constrained isostere of ciliobrevin recently described *in vitro* to perturb retrograde transport.³³

The comprehensive measurements of translational and rotational motions in these models provided by our nonlinear nanoparticle-based assay will allow to finely study axonal transport in the central nervous system *in vivo*. In particular, our method may be instrumental to investigate the biological significance of altered cargo pausing duration and run length and to study the link between transport defects and neuronal dysfunctions. In this context, the zebrafish model will be amenable to conduct high-content screening of therapeutic drugs able to rescue axonal transport impairment.

Results and Discussion

***In vivo* measurement of intraneuronal transport.** To investigate the axonal transport in the CNS of Zf larvae imaged by fast nonlinear microscopy, we synthesized single crystal nanoKTP (Supporting Figure S1A) in solution in deionised water as we reported previously.²⁶ Using differential centrifugation we narrowed their size distribution, measured by nanoparticle tracking analysis, leading to an average size of 123.8 ± 2.4 nm, with a standard deviation of 42.7 nm (Supporting Data S1 and Figure S1B). This solution was injected in the left optic tectum of 3 days post-fertilization (dpf) larvae at a concentration of ≈ 1 mg/mL (Fig. 1A). We conducted the nonlinear microscopy fast-scanning recording 24 hours later, at 4 dpf, when tectal neurons are mature and respond to visual stimulations.³⁴ Prior to live imaging of the optic tectum, Zf larvae were anesthetized and placed in dorsal view in an agar mold (Fig. 1B). Once in the right orientation, larvae were immobilized by covering them with a cushion of low-melting agarose maintained during the acquisition, performed at 24°C. The injection condition (solution concentration and volume) were optimized to obtain a sparse labeling, but sufficiently dense to collect enough data to ensure reliable statistical analysis in order to minimize the risk of interference with the endocytosis along the axons, which is a key physiological process for neuronal polarity maintenance.³⁵ Failure of the injection was rare: about 90% of injected larvae contained enough nanoKTP in the region of interest for axonal transport analysis.

We set the excitation laser wavelength at 1040 nm, and its power at 12.5 mW at the exit of the microscope objective ($\times 25$, water immersion, numerical aperture 0.95). This power value was chosen to take advantage of the full dynamic range of the detector (8-bits

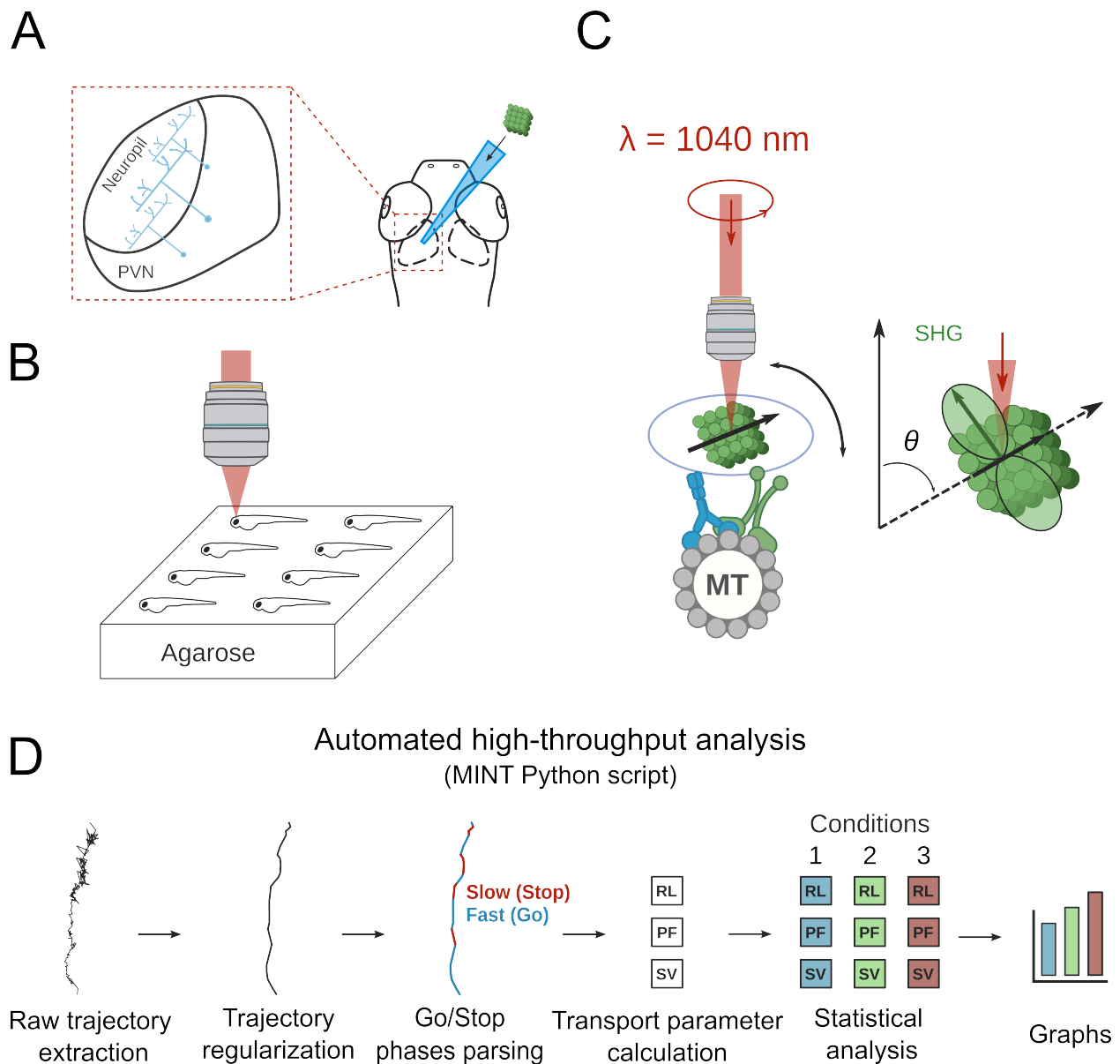


Figure 1: Method to measure intraneuronal transport parameters in the brain of zebrafish larvae, based on nonlinear nanocrystals microinjection and endocytosis, second-harmonic generation fast microscopy, and high-throughput automatic extraction and analysis of nanocrystal-labeled vesicle trajectories. (Continued on the following page).

encoding, maximum pixel intensity of 255). As the nanoKTP SHG is mainly excited by an electric-field direction along a single crystalline axis we use a circularly polarized excitation beam (Fig. 1C) in order to favor the excitation of all the particles whatever their orientation. The light from the nanoKTP is collected with the same microscope objective, then passed through a bandpass filter (50 nm spectral width) centered on 525 nm wavelength, before being detected by a point detector, without any polarization selection. Taking advantage of prism-based tunable emission spectrum detection of our microscope, we recorded the signal

Figure 1: (A) Microinjection of the nanoKTP (green cube) in the left optic tectum of the head of a zebrafish larva (at 3 dpf). Inset: zoom on optic tectum displaying periventricular neurons (PVN) projecting parallel and coplanar axons in the neuropil. (B) Immobilization at 4 dpf of 8 anesthetized injected larvae in channels molded in agarose, with dorsal orientation towards the microscope objective for SHG microscopy. (C) Left: schematics of a single monocrystalline nanoKTP embedded in a vesicle connected to a microtubule (displayed as a section, MT) by a dynein (blue) and a kinesin (green). The black arrow onto the nanoKTP indicates the direction of the second order nonlinear polarization (mostly dipolar), in response to the pulsed-laser excitation at 1040 nm wavelength. Right: in green, the SHG radiating pattern of one nanoKTP. (D) Analysis pipeline with the Python program MINT we developed,³⁶ consisting, from left to right, in two parts of analysis. The first part includes (i) trajectories extraction, (ii) trajectory regularization (filtering out nanoparticle position noise) and (iii) parsing in fast (“Go”) and slow/pause (“Stop”) phases of motion. The second part starts with (i) the calculation of transport parameters for each trajectory, among which the segmental velocity (SV), which is the average velocity during a Go phase segment, the run length (RL), which is the distance traveled between two consecutive Stops, the pausing frequency (PF), etc and continuing with (ii) the analysis of the data for different biological conditions provided as inputs, transport parameters statistical comparison, and finally (iii) the plotting of graphs associated to these data.

emission spectrum from a single nanoKTP and confirmed that it corresponds solely to SHG signal (Supporting Data S2 and Figure S2).

Driving the mirror galvanometers of the microscope at their resonance frequency (8 kHz), we acquired continuous 512×352 pixels raster scans at a constant depth position ($\approx 100 \mu\text{m}$) below the skin with a line average of 2, leading to a refreshing rate of 20 frames/s (*i.e.* 50 ms per full frame scan) identical to the one we use in wide-field acquisition on cultured neurons,^{22,37} that enhances the sensitivity to short pauses. In order to avoid degradation of the diffraction limited spatial resolution by sampling, we selected a pixel size of 173 nm, which fulfills the Shannon criteria as it is smaller than half the diffraction-limited spot radius of two-photon-excited microscopy $\rho_{\text{Airy}}^2 \text{ph}/2$. The Airy radius in two-photon-excited process can indeed be estimated by $\rho_{\text{Airy}}^2 \text{ph} \equiv 1.22\lambda/(2\sqrt{2}NA)$, leading to $\rho_{\text{Airy}}^2 \text{ph}/2 \approx 236 \text{ nm} > 173 \text{ nm}$, at the excitation wavelength of $\lambda = 1040 \text{ nm}$ and for $NA = 0.95$ microscope objective numerical aperture.

With this pixel size value of 173 nm, the scan covers $88.6 \mu\text{m} \times 60.9 \mu\text{m}$. We set the total number of frames in the video to either 2354 (Supporting Video S1) or 2501 (Supporting Video S2), which corresponds to video durations of about 120 s. Compared to the wide-field mode, for which the integration time per pixel is 50 ms, here in the raster-scan mode, the duration of photon counting (dwell time) is only 144 ns/pixel (taking into account the 2-lines averaging done while scanning). To collect a sufficient number of trajectories per larvae, we record 4 videos corresponding to different regions of interest (and different depths) in the optic tectum ipsi- or contra-lateral to the injection site (total of 8 videos per larva). We did not observe mortality due to intracerebral injections or imaging. The acquired videos are then processed by an automated high-throughput analysis pipeline (Fig. 1D) detailed later.

Internalisation and distribution of the nanoKTP in the periventricular neurons of the CNS.

Figure 2A displays the average projection in time of Supporting Video 1. It shows that nanoKTP motions appear mostly parallel and oriented radially relative to the eye (top left corner), as expected in this brain region (Fig. 1A, inset). This suggests their internalization in neurons, whose projections match the axons of PVN. Here we stress that this endolysosomal vesicle labeling strategy (via nanoKTP spontaneous uptake) is naturally sparse and does not require further sorting.¹¹⁻¹³ To better identify the cell type that was targeted by the nanoKTP, we performed simultaneous live labeling of nanoKTP with the cell membrane dye DiI. The pattern of expression obtained in two-photon-excited fluorescence microscopy reveals DiI positive cells with the morphology of PVN and the presence of nanoKTP in the axon (Fig. 2B). To confirm these observations, we conducted immunofluorescence and quantified the localization of the nanoKTP in different cell subtypes present at the site of injection. Twenty four hours post injection, Zf larvae were immunostained to label acetylated-tubulin (ac-tub), considered as a marker of axons, and glial fibrillary acidic protein (GFAP), as a marker of glial cells and neuronal precursors (Fig. 2C-E). Nanoparticles colocalize in 27% of the cases with ac-tub-positive axons, which is 2.7 times more than the colocalization with GFAP-positive branches presenting 10.2% of colocalization (Fig. 2F). In about 60% of the cases, the nanoKTP were found aside from ac-tub and GFAP-positive structures, which might correspond to particles stuck in the extracellular matrix, unable to diffuse to axonal membranes where they may be endocytosed.

As all PVN bodies are gathered in the PVN-labeled area of Fig. 1A inset (between the neuropil and the ventricle) and project towards the neuropil (located above the eye), motions towards negative values of x (Ox axis pointing right, with origin in the eye, see Fig. 2A), are anterograde if we are addressing the left eye, and retrograde for the right eye.

Note that in Fig. 2A, most lines are made of a succession of bright and dim spots. This blinking reflects the rotational motion (relative to the microscope objective axis (Oz)) of the vesicle-containing nanoKTP, upon their transport by the molecular motors stepping on the microtubules. This intensity fluctuation is a consequence of the anisotropy of nanoKTP SHG radiation pattern that can be assimilated to the one of a dipole (Fig. 1C). Indeed, KTP second-order nonlinear susceptibility matrix³⁸ has a dominant coefficient leading to a dipolar nonlinear polarization mostly parallel to the c crystallographic axis. When the dipole is aligned with the objective axis, almost no light is collected, leading to a very dim signal. Incidentally, the large variation of intensity of the vast majority of spots indicate that each endosome most probably contain a single nanoKTP, because the presence, within a vesicle, of multiple randomly oriented nanocrystals would yield a much lower intensity contrast. This observation confirms that *in vivo*, like in primary cultures,^{22,37} neuron internalize at most one solid nanoparticle per vesicle. The nanoKTP still appear as single particles per transported vesicular compartment 24 hours after injection, despite the possibility for endolysosomes to merge. This is probably due to the sparse labeling we use: only a small proportion of vesicles are labeled so it is highly unlikely that two endolysosomes destined for fusion would both contain a nanoKTP.

In the subsequent studies, we evaluated the fraction of moving particles in large sets of data and found $6.7 \pm 0.6\%$ (standard error on the mean, s.e.m.) (Fig. 4B) and $8.6 \pm 0.8\%$ (Fig. 5B) in two independent studies, leading to an average of 7.7%. We can conclude that, statistically, a non-negligible fraction of $7.7/27 \approx 28\%$ of the nanoKTP colocalizing with

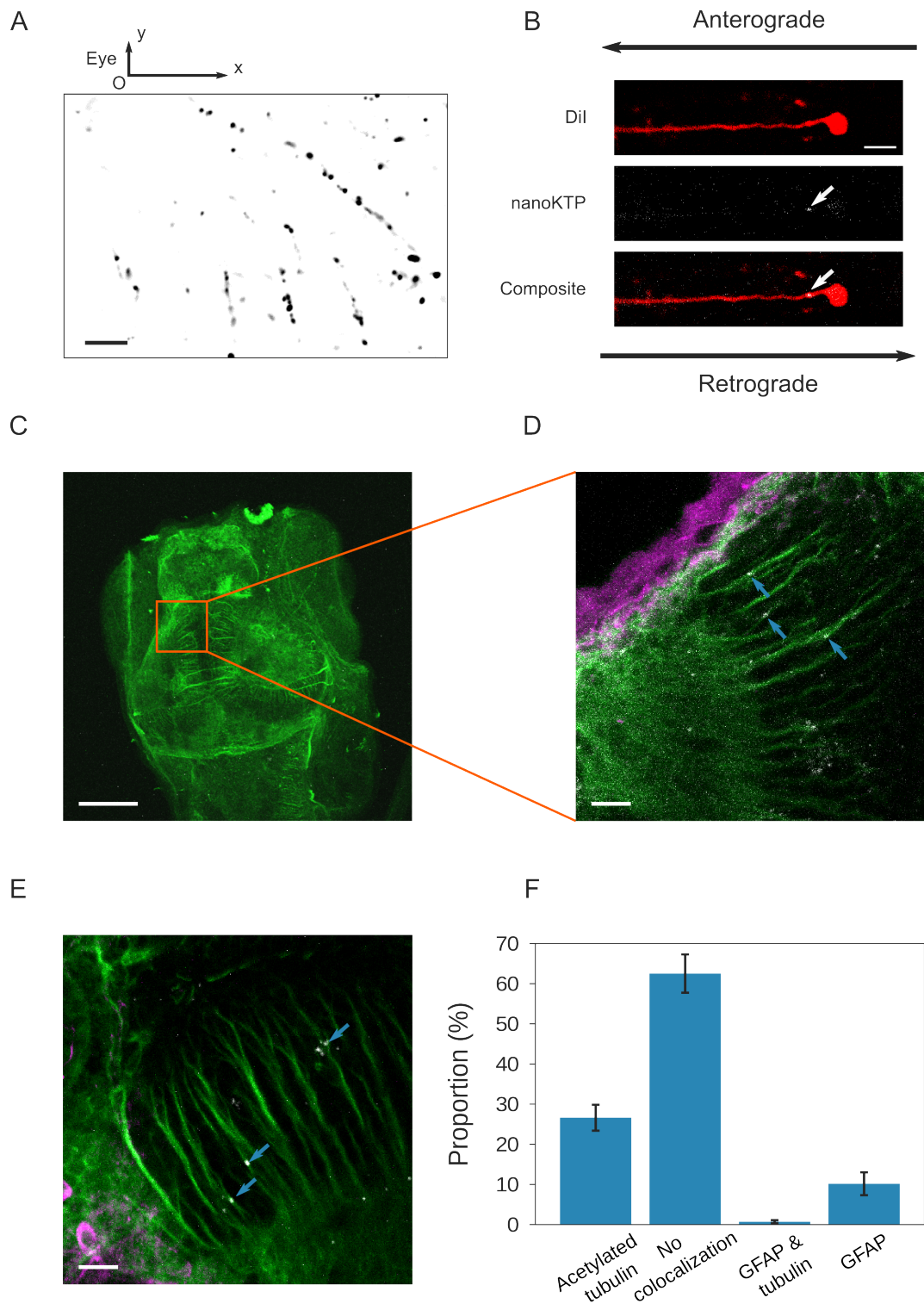


Figure 2: Localization of the nanoKTP relative to axons and glial cells of the zebrafish larva optic tectum injection zone. (Continued on the following page).

axons, moves during the 2-minutes raster-scans recording. This value must be compared to the mobile fractions of $\approx 60\%$ reported for various endosomes in motor neurons of Zf

Figure 2: (A) Average intensity projection of a single plane raster-scan video (Supporting Video 1) of the SHG signal from nanoKTP in the brain of a Zf larvae (at 4 dpf). Moving nanoKTP have directed, mostly linear, displacements oriented towards the left top corner where the larva left eye is located (out of the field of view, FoV). Scale bar: 10 μm . (B) Top: two-photon fluorescence image of a DiI cyanine dye-labeled periventricular neuron in a live Zf (4 dpf); middle: SHG signal of the same FoV as the top panel, showing in white a nanoKTP (white arrow) colocalized with the axon of the PVN. Bottom: merge of top and middle images. Retrograde and anterograde directions are indicated by arrows. Scale bar: 10 μm . (C) Acetylated tubulin immunofluorescence labeling of a zebrafish larva head (in ventral position). Maximum projection of confocal slices of a z -stack covering a thickness of 30 μm . Scale bar: 100 μm . (D) Representative field of view (zoom of (C) larva) merging three detection channels: acetylated tubulin (ac-tub, green) and GFAP (magenta) immunostainings, and nanoKTP (white). Maximal projection over a total thickness of 5.1 μm . Nanoparticles colocalized with ac-tub are pointed by arrows. (E) Another FoV with the same labels as in (D). Maximal projection over a total thickness of 4 μm . Scale bars for (D) and (E): 10 μm . (F) Proportion of the localization of nanoKTP relative to ac-tub and GFAP immunostaining ($n = 4$ larvae and 2-7 FoV per larva). Error bars: s.e.m.

larvae spinal cord.¹² Beyond the fact that our study addresses axonal transport in CNS neurons, that may have different dynamics compared to motor neurons, but also different endolysosomal compartments than in Ref.,¹² the 2-fold discrepancy observed may be due to an overestimation of nanoKTP internalized in axons. Indeed diffraction limited imaging of sub-micrometer sized structures like axons and nanoKTP makes it difficult to discriminate between particles that stayed at axonal outer-membrane from the ones that were fully internalized.

Key parameters for quantitative analysis of axonal transport with nanoKTP.

Imaging data sets were further analyzed to extract quantitative parameters required to assess axonal transport defects. We first evaluated the precision of localization and applied a denoising strategy before processing data through the automated pipeline of transport analysis schematized on Fig. 1D. Trajectories of nanoKTP-labeled vesicles are inferred from consecutive positions of the nanoparticles (Materials and Methods) obtained as the position of the center of the best Gaussian fit to each SHG spot. Focusing on directed motions, we first filter-out “trajectories” corresponding to confined Brownian motions (possibly from particles in intercellular space) that are characterized by a bounded mean-square displacement.³⁹

In the remaining trajectories, the detected positions are affected by a precision of localization noise whose 2D-standard deviation σ_{xy} is expected to scale down with the number of detected photons N , as $\sigma_{xy} \simeq s/\sqrt{N}$, in the case of a photon shot noise-limited process, where s is the standard deviation of the microscope objective experimental point spread function (PSF_{exp}). We estimated N after having determined a conversion factor between the 8-bits grey level of the hybrid photomultiplier detector (Supporting Data S3 and Figure S3), and we inferred $s \approx 355$ nm from the smallest SHG spots (Supporting Data S4 and Figure S4A).

We measured $\sigma_{xy}(N)$ from videos of static nanoKTP (dropcasted on a glass coverslip) in the same acquisition conditions as for the intraneuronal transport (20 fps, in the cage incubator). Supporting Figure S4A displays one SHG spot from Supporting Video S3 and Gaussian fits of two orthogonal cross-sections of this spot. Figure S4B is a scatter plot of Gaussian fit centers positions of the same nanoKTP spot in consecutive frames. The precision of localization of this spot is given by the $\sigma_{xy} \equiv \sqrt{\sigma_x^2 + \sigma_y^2}$, where σ_x and σ_y are the standard deviations in x and y directions.

We then considered an ensemble of SHG spots with different N and plotted on Supporting Figure S4C $\sigma_{xy}(N)$. The smallest experimental precision is $\sigma_{xy}^{\text{min,exp}} \approx 50$ nm, obtained for $N = 425$ photons, while the theoretical minimal value for the same N is 10 nm (Supporting Data S4). We attribute this ≈ 40 nm excess of noise mostly to mechanical vibrations. At the other extremity, the worst localization precision of $\sigma_{xy}^{\text{max,exp}} = 371$ nm correspond to the dimmest spot. Altogether the average value is $\bar{\sigma}_{xy}^{\text{exp}} = 129$ nm.

Considering the blinking of the SHG signal due to the vesicle rotational mobility, a spot that appears bright in one frame and can be localized with 50 nm precision, may also become very dim a few frames after and be localized with a precision of only ≈ 370 nm. Such variations along one trajectory may induce artifacts in transport parameters quantification. For instance, in the extreme case of two consecutive positions separated by 370 nm, we cannot discriminate between a static faint particle and a very fast displacement, at an instantaneous velocity of $0.370/0.05 \approx 7.4$ $\mu\text{m/s}$, which is very unlikely as the average reported velocity in zebrafish neurons is in the 0.4-1.2 $\mu\text{m/s}$ range.⁴⁰

In order to mitigate the impact on transport parameters values of a low SHG intensity (associated to a large localization uncertainty), we regularized the trajectories by a nonlinear total variation-based noise removal algorithm⁴¹ (Fig. 1D). As the localization uncertainty introduces variations of the instantaneous velocity larger than the one expected from the true vesicular motion (constituted of phases of almost constant velocity, sparsely interrupted by much slower motion or complete pauses) we implemented a regularization algorithm (see Materials and Methods) that infer the positions' ground truth by minimizing the acceleration (velocity changes), under a constraint related to the average experimental precision of localization of $\bar{\sigma}_{xy}^{\text{exp}}$ previously determined separately.

Based on the straight pattern of PVN axonal projections, endolysosomal trajectories are expected not to deviate strongly from straight lines. Hence, at this stage of the analysis, we excluded the trajectories that were not properly fitted by a third order polynomial law (Materials and Methods), which represented 17% of them.

The following step of the automated analysis consists in parsing the regularized trajectories in phases of "slow" (further called Stop) and "fast" (Go) motions (Fig. 1D). To this aim we apply the same heuristic method as in our previous work.²² Briefly, we calculate at each trajectory position i a confinement ratio R_i on a sliding window of 6 consecutive points width, defined as the net distance between the first and last point divided by the sum of the length of the two segments. $R_i = 1$ corresponds to straight line, and therefore a Go phase, while $R_i = 0$ is associated to a zero net motion, hence a Stop phase. To attribute a phase state to all other intermediate values of R_i , we searched for a confinement ratio threshold R_{th} emerging from the datasets. To this aim we examined the distribution of R_i for all trajectories of all conditions (including heterozygote larvae) in *kif5aa* larvae (Supporting Data S5 and Figure S5) and noticed that it has a minimum at 0.64. We then decided to consider this

minimum as the threshold to parse the phase states: i^{th} point with $R_i > R_{\text{th}} = 0.64$ is in a Go phase, while for $R_i \leq R_{\text{th}}$ the vesicle is in a Stop phase.

For the quantitative analysis of the transport, we calculate two general parameters and six metrics associated to individual trajectories, distinguishing retrograde from anterograde displacements according to previously described criteria (*i.e.* the sign of the displacement along (Ox) and the eye side). The two general parameters are the fraction of moving particles detected in each video (*i.e.* in each field of view), and the transport directionality evaluated as the proportions of purely retrograde (R), anterograde (A) or bidirectional trajectories (B). In the case of bidirectional trajectories, we refine the analysis by calculating the fraction of retrograde Go phase over all types of phases (calculated from the traveled distances).

We then characterize each trajectory by the six metrics inferred from the parsing: retrograde and anterograde segmental velocities (SV, average velocities during R and A Go phases only), the pausing frequency (PF, number of Stop phases per minutes), the average pausing duration (PD), the retrograde and anterograde run lengths (RL, average distances traveled during R and A phases).

In order to check that the mere injection of nanoKTP does not impact axonal transport, we investigated lysosomal transport, in wild type and injected conditions. This was assessed by incubating the larvae with LysoTracker Red, a live labeling dye of lysosomes. To quantify the axonal transport of LysoTracker-positive vesicles, we used exactly the same pipeline of recording and analysis than for nanoKTP, here taking advantage of the two-photon absorption of the dye. Despite a lower signal-to-background ratio for LysoTracker than for nanoKTP and bleaching, we were able to extract enough reliable trajectories to compare the transport parameters with or without nanoKTP, and did not find any difference between these two conditions as shown in Supporting Figure S6. Hence our method, with its sparse labeling of tracked compartments (as shown on Fig. 2E-F), does not only have no measurable impact on lysosomal transport parameters, but it also provides a larger number of reliable trajectories, owing to the large signal-to-background and perfect photostability of the SHG signal.

This LysoTracker Red labeling experiment gave us also the opportunity to evaluate the colocalization of nanoKTP with late endosomes or lysosomes, which are the endolysosomal compartments positive to LysoTracker. We only considered live colocalizations of nanoKTP having directed motion and found that about 56% of them colocalize with LysoTracker Red, as shown in Supporting Figure S7. This observation is consistent with the timeline of the endocytosis cycle, since we record nanoKTP motion 24h after their injection, at a stage where the majority of the up-taken nanoparticles is expected to have reached late endosomes and lysosomes. The remaining fraction of 44% may correspond to nanoKTP that are in less acidic endosomal compartments, with lower LysoTracker Red signal. Hereafter, we name the mobile nanoKTP containing compartments “endolysosomal vesicles”, shortened sometimes in “vesicles”.

In the following we show that the general parameters and trajectory metrics defined above and evaluated for nanoKTP-labeled endolysosomal vesicles, are sensitive enough to measure the impact of drugs inhibiting the motors, or loss-of-function mutations, starting with dynein (using either a mutant of dynein heavy chain 1 or a dynein inhibitor) and continuing with the kinesin 1 mutant *kif5aa*.

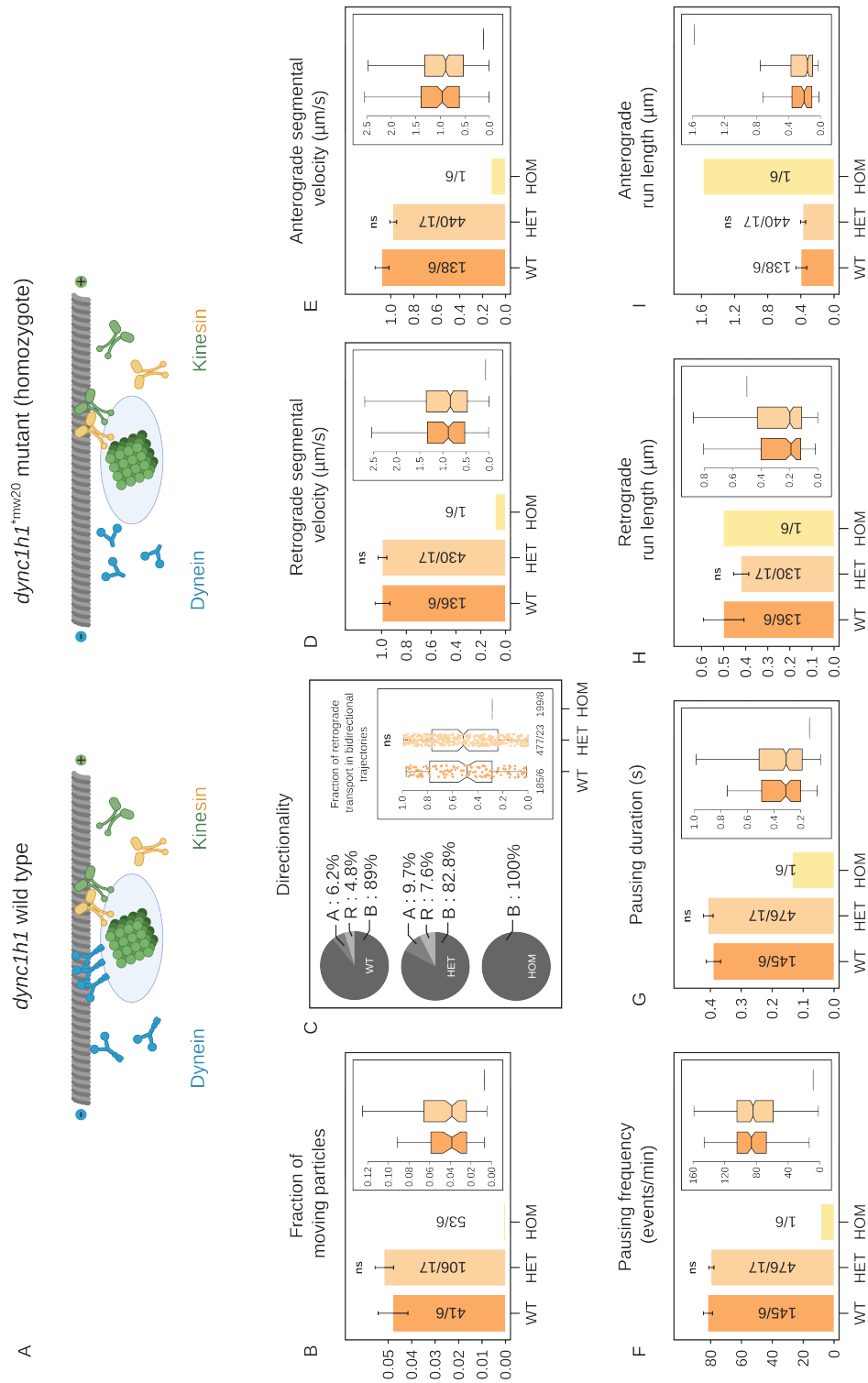


Figure 3: *dync1h1*^{mw20} dynein mutation stops intraneuronal transport in zebrafish brain. (Continued on the following page).

Mutation or inhibition of dynein motors affects endolysosomal transport *in vivo*. We first consider the *mw20* zebrafish transgenic line with a point mutation of *dync1h1*

Figure 3: (A) Schematic representation of the mechanisms of intraneuronal transport. Left: Typical transport conditions in a wild type *dync1h1^{mw20}* sibling. The optically active nanoparticle (green cube) is spontaneously internalized in an endolysosomal vesicle (light blue oval) and is then transported along a microtubule (grey tube) in an anterograde motion by kinesins (green and yellow dimers to represent two different families, including kinesin 1) or in a retrograde motion by dyneins (blue dimers). Right: Impaired transport conditions by mutant dynein heavy chain (blue shortened dimers). (B–I) Modifications of intraneuronal transport parameters in *dync1h1^{mw20}* larvae brain, depending on the larva genotype (WT: wild type, HET: heterozygote and HOM: homozygote). For (B, D–I) bar plots display the parameter mean values \pm s.e.m., while insets are distribution box plots of the same parameter. In (B) numbers in the bars are # of fields of view analyzed/# of animals analyzed, while for (C–I) they are # of trajectories/# of animals analyzed. (B) Fraction of moving particles, with directed motion. The 3-conditions Kruskal-Wallis p -value=0.22. The mobile fraction in HOM mutant is 0.01% while it is 4.8% and 5.2% in WT and HET respectively. (C) Directionality. Left : pie charts of the percentage of purely anterograde [A], purely retrograde [R], or bidirectional trajectories [B], for each condition. Right inset: fraction of R transport in bidirectional trajectories. (i.e., having at least one phase of R and one of A transport). A value of 1 means a purely retrograde trajectory, while a value of 0 signifies a purely anterograde one. No difference are observed ($p = 0.74$). (D–E) Retrograde and anterograde segmental velocities do not vary between WT and HET ($p = 0.84$ and $p = 0.26$). Overall Kruskal-Wallis test p -value =0.17 for A and p -value =0.29 for R. The A and R phases in the only bidirectional trajectory have very slow segmental velocities of $0.07 \mu\text{m/s}$ for R and $0.12 \mu\text{m/s}$ for A. (F) Pausing frequency. Overall Kruskal-Wallis p -value = 0.23. PF does not vary between WT and HET ($p = 0.49$). In HOM the pausing frequency is ten times smaller than in WT and HET (8.2 events/s compared ≈ 80 events/s) (G) Pausing duration does not differs between conditions ($p = 0.347$). There is no difference between WT and HET ($p = 0.836$) but the pausing duration is 3-fold smaller in the trajectory of the HOM mutant. (H) Retrograde run length does not differ between conditions ($p = 0.572$). (I) Anterograde run length does not vary across the different conditions ($p = 0.21$). The RL of $\approx 1.6 \mu\text{m}$ for the only HOM trajectory is not representative. RL values of $\approx 0.4 \mu\text{m}$ for WT and HET are consistent with the one reported for *kif5aa* and DYNA data.

(*dync1h1^{mw20}*) coding for heavy chain 1 of the dynein motor complex. This mutation creates a premature stop codon leading to a non-functional dynein devoid of the tail (that binds to the microtubule, see Figure 3A) and of the domains associated to ATPase activity.³¹ Despite this mutation, the maternal contribution of wild-type *dync1h1* mRNA and protein leads to larvae survival at least during 8 days, which fits our experimental plan.

Fig. 3(B–I) display axonal transport properties in homozygous (HOM) mutant, heterozygous (HET) or wild-type (WT) siblings. We use the fraction of moving particles in each field of view (of similar morphology) as a first parameter to quantify the overall impact of the mutation on the transport. In homozygous *dync1h1^{mw20}* mutant data set we detected only one particle having a directed motion corresponding to an estimated mobile fraction of 0.01%, in contrast to the wild-type and heterozygous larvae in which this fraction respectively reached

4.8% and 5.2% (Fig. 3B). Furthermore, we observed no differences between WT and HET larvae in any of the other transport parameters (Fig. 3(C-I)). We attribute these results to the fact that most dynein motor complexes are inactive even in normal conditions as they need to be associated to activators (as shown in Redwine *et al.*⁴²), making it possible to maintain a normal transport with only half of the WT concentration of dynein heavy chain 1, as long as this half forms the same concentration of active complexes than in WT conditions. Such a phenomenon may explain the absence of transport parameter modifications in HET *dync1h1*^{mw20} larvae compared to WT siblings. Regarding the vesicle directionality (pie plots of Fig. 3C), the dominant category of bidirectional motion within single trajectories is consistent with observations reported in experiments using high-resolution analysis like ours,²³ and most likely reveals a tug-of-war situation of motors pulling in opposite direction alternatively, leading however to a net displacement which is either anterograde or retrograde. As a refinement, we also investigated the fraction of retrograde transport within the population of bidirectional trajectories only (inset of Figure 3C). This fraction would be equal to 1 for a trajectory made of purely retrograde segments. It did not show differences between WT and HET.

Dynein being the only retrograde motor, a strongly modified retrograde transport was expected in *dync1h1*^{mw20} mutants, but the anterograde transport was also disrupted. Indeed, this observation is consistent with several reports in which the interruption of retrograde transport, in particular *via* a mutation on dynactin (complexing with dynein to activate it), impacts the anterograde transport too.⁴³⁻⁴⁵

As the endolysosomal transport phenotype of *dync1h1*^{mw20} homozygous larvae is severe, we decided to challenge our axonal transport measurement method with another model of retrograde transport impairment. Zebrafish larvae were then treated with a dynein inhibitor for which concentration can be adjusted to produce moderate effects. To this aim, we selected dynapyrazole-A (DYNA), a new conformational isostere of the first-in-class dynein inhibitor ciliobrevin, that has a higher selectivity in blocking only dynein's microtubule-stimulated activity (Figure 4A) and not the ATPase one.³³

In *in vitro* assays DYNA strongly inhibits dynein's microtubule-stimulated activity at micromolar concentrations. In absence of data available *in vivo*, we first immersed Zf larvae in different concentration (10 and 15 μ M) of dynapyrazole for 25 to 45 min at 28°C in a bath. The larvae were then washed and observed to monitor the appearance of potential toxicity effects right after or 1 h post exposure. Thirty minutes incubation at 15 μ M lead to severe malformations (body curvature) detected 1 h post exposure, which evolved into larvae death (data not shown). As there was no toxicity at 10 μ M (25 min exposure), the following experiments were performed at this concentration. Figure 4B-I display the effect of DYNA on the axonal transport properties. In untreated control sample, the fraction of moving particles in each field of view is about 6.7%. while it decreases to 4.6% upon DYNA exposure, representing a 32% decrease (Figure 4B). As DYNA is not known to impact endocytosis, we attribute this reduction to the inhibition of dynein's microtubule-stimulated activity. Our observations are in agreement with the transport impairment of fluorescently labeled lysosomes initially reported *in vitro*.³³ We then questioned whether this dynein inhibition modifies the preferred direction of transport. Pie plots of Figure 4C displays the proportions of purely retrograde, anterograde or bidirectional trajectories for both control and DYNA conditions. The total fraction of purely directional trajectories is slightly reduced by DYNA

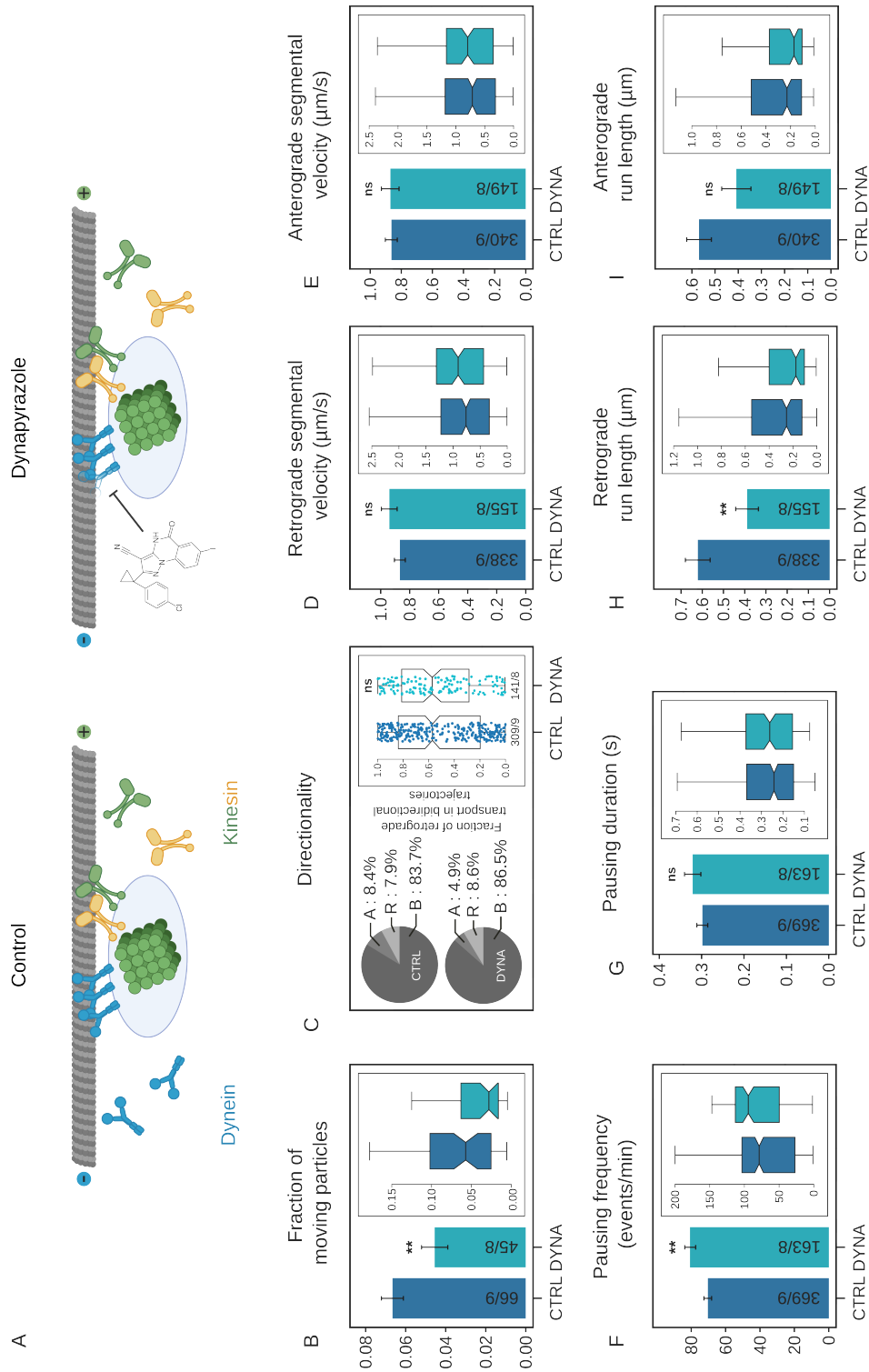


Figure 4: Dynein inhibitor dynapyrazole impairs intraneuronal transport in zebrafish brain. (Continued on the following page).

(16.3% in control to 13.5% in DYNA). In a counter-intuitive way, within this fraction, the one of purely anterograde transport decreases by 42% from 8.4% (control) to 4.9% (DYNA),

Figure 4: (A) Schematic representation of the mechanisms of intraneuronal transport. Left: Typical transport conditions in a control animal. NanoKTP (green cube) is spontaneously endocytosed and 24h after, ends up in an endolysosomal vesicle (light blue oval) of a PVN axon and is transported along a microtubule (MT, grey tube) in the anterograde direction (towards MT + end) by kinesins (green and yellow) or in the retrograde direction (towards MT – end) by dyneins (blue). Right : dynapyrazole impairment of microtubule-stimulated dynein activity. (B-I) Modifications of transport parameters in larvae bathed in dynapyrazole at 10 μ M concentration (condition DYNA), compared to control (bath without dynapyrazole, CTRL). For (B, D-I) bar plots display the parameter mean values \pm s.e.m., while insets are distribution box plots of the same parameters. In (B) numbers in the bars are # of fields of view analyzed/# of animals analyzed, while for (C-I) they are # of trajectories/# of animals analyzed. (B) Fraction of moving particles, with directed motion. The treated larvae have 31% less moving particles than the control ($p = 0.004$). (C) Transport directionality. Pie charts of the percentage of purely anterograde (labeled A), purely retrograde (R), or bidirectional trajectories (B), for each condition. The proportion of purely A trajectories decreases in favor of purely R trajectories in treated larvae. Inset : fraction of retrograde transport in bidirectional trajectories. This fraction does not vary significantly ($p = 0.738$). (D-E) Retrograde and anterograde segmental velocities. Both R and A segmental velocities are not statistically different between the different conditions ($p = 0.161$ and 0.885 , respectively). Trajectories with purely A and purely R movement were excluded from the analysis of R and A segmental velocities, respectively. (F) Pausing frequency. The pausing frequency of the treated larvae is 14% larger than that of the control ($p = 0.004$). (G) Pausing duration does not change under DYNA treatment ($p = 0.261$). (H) Retrograde run length is 37% shorter for the treated larvae ($p = 0.004$) compared to the control. Trajectories with purely anterograde movement were excluded from analysis. (I) Anterograde run length decreases by 28% in treated larvae, but there are no statistical differences between conditions ($p = 0.084$). Trajectories with purely retrograde movement were excluded from analysis.

while purely retrograde transport stays almost constant evolving from 7.9% (control) to 8.6% (DYNA). The fraction of retrograde transport within the population of bidirectional trajectories only (inset of Figure 4C) has a very broad distribution, indicating that even bidirectional trajectories are polarized (with a slight bias towards R direction as evidenced by the value of the median of 0.57), leading to a net displacement in either R or A direction. We observed that the fraction of retrograde transport in these bidirectional trajectories does not vary significantly between control and DYNA, but we noticed however that its distribution slightly narrows from control to DYNA, which is the sign of a slight decrease of polarity. This conclusion is further confirmed by the significant increase of 19% of the number of direction reversals per minute within a trajectory (Supporting Figure S8A).

We then focused our attention on specific parameters of the retrograde transport. We did not detect significant variations of the R and A segmental velocities (Figure 4D-E) which values are around $0.9 \mu\text{m/s}$ for both directions, consistent with values reported for axonal transport of various endosomes in Zf larvae motor neurons.^{12,40} On the contrary we observed a significant increase of about 15% of the pausing frequency (Figure 4F) with DYNA (from

70.4 to 80.7 events/min), accompanied with a trend of augmentation of the pausing duration, all types of pauses included (Figure 4G, from 0.30 to 0.32 s). We investigated whether this trend could be due to a sub-population of pauses, considering the ones between either anterograde, opposite directions, of retrograde phase of motions (Supporting Figure S9A-C). None of these categories revealed any differences of pausing duration for DYNA treated Zf compared to control.

Regarding the spatial parameters, we measured a significant decrease of the retrograde run length by 37% from 622 to 390 nm (Figure 4H) concomitant with a trend of decrease by 28% of anterograde run length from 570 to 409 nm (Figure 4I). Our observations are in agreement with previous reports in which the reduction of cytoplasmic dynein concentration was observed to impair the transport parameters⁴⁶ with a similar reduction of anterograde RL, indicating that dynein is also involved in the activation of anterograde transport.

We then considered the perturbation of motors driving the anterograde transport in PVN neurons of Zf brain. To this aim we investigated the effect of a change of concentration of the dominant kinesin motor in the brain, kinesin 1, as induced in a mutant zebrafish.

Precise measurement of slight endolysosomal transport defects induced by a kinesin 1 motor disruption. Anterograde vesicle displacement along microtubule is mediated by kinesin superfamily proteins⁴ that also play a key role in the organism development.⁴⁷ To validate our model of *in vivo* detection of fine axonal transport impairment caused by abnormal kinesin concentration, we monitored nanoKTP axonal transport in the *kif5aa* zebrafish transgenic line. This mutant generated by Auer *et al.*,³² was engineered to introduce a premature stop in the motor part of Kif5aa molecule, hence preventing its complete synthesis and therefore its function, leading to abnormal phenotype that includes blindness and failure to inflate the swim bladder, eventually resulting in death around 10 dpf.

Moreover, this mutant is relevant to our experimental demonstration as *kif5aa* expression was shown to be specific to neurons,⁴⁸ and in wild-type siblings of *kif5aa* transgenic line at 3 dpf, *kif5aa* is expressed in the whole optic tectum, including PVN neurons (refer to Fig.1B of ref.³²). *Kif5aa* mRNA is also down-regulated in mutant larvae probably due to nonsense-mediated decay of the mutant transcript containing a premature stop codon that leads to the translation of a non functional truncated protein. Considering also that *kif5aa* is not maternally expressed in the germline,⁴⁸ and therefore Kif5aa not maternally contributed, we expect that, in *kif5aa* mutant, the level of functional Kif5aa in axons of PVN (the brain region we studied) is smaller than in wild-type siblings. However, other types of kinesins could compensate the decrease of functional Kif5aa and normalize the transport, including the protein products of the other four *kif5* paralogs present in Zf encoding for kinesin 1 heavy chain isoforms.⁴⁸ This is in contrast to the previous situation where dynein represents the only motor driving retrograde transport.

In *kif5aa* larvae (Figure 5A) we evaluated the same axonal transport parameters as in dynein inhibition experiment, comparing mutant (MUT) to wild-type siblings (WT). We did not detect differences in the fraction of moving particles between these two conditions in axons of the PVN (Figure 5B). These observations differ from previously published results conducted in the axons of retinal ganglion cells of *kif5aa* mutants showing an increase of the mobile fraction of small vesicles.³²

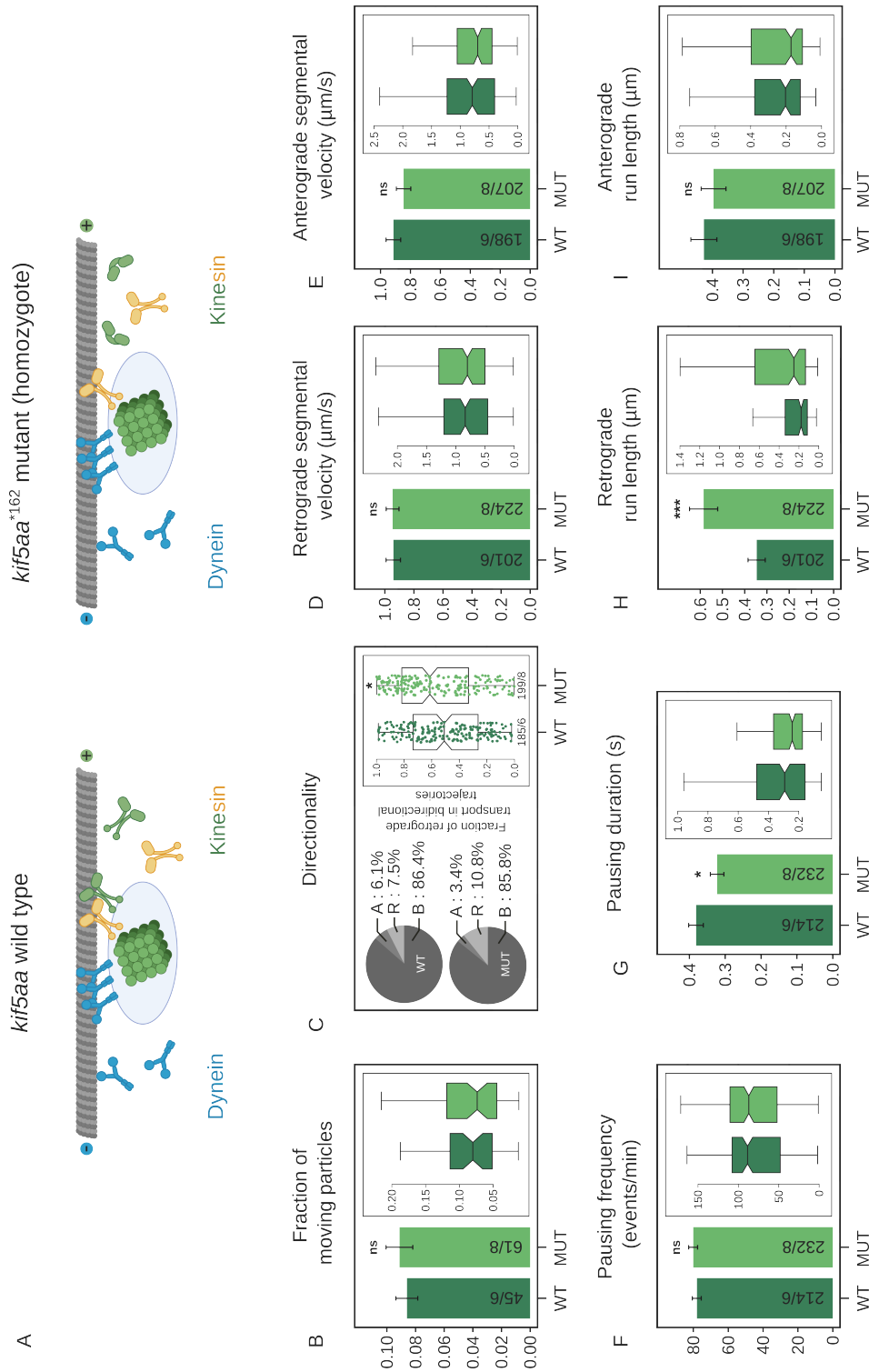


Figure 5: *kif5aa* mutation impairs intraneuronal transport in zebrafish brain. (Continued on the following page).

Then, we investigated the transport directionality (pie plots of Figure 5C). In mutant larvae, while the total fraction of purely directional motion only slightly increases from 13.6%

Figure 5: (A) Schematic representation of the mechanisms of intraneuronal transport. Left: Typical transport conditions in a wild type *kif5aa* sibling. The optically active nanoparticle (green cube) is spontaneously internalized in an endolysosomal vesicle (light blue oval) and is then transported along a microtubule (grey tube) in an anterograde motion by kinesins (green and yellow) or in a retrograde motion by dyneins (blue). Right: Impaired transport conditions by mutant kinesins (green shortened dimers). (B–I) Modifications of intraneuronal transport parameters in *kif5aa* larvae brain, depending on the larva genotype (WT: wild type or MUT: mutant homozygote). For (B, D-I) bar plots display the parameter mean values \pm s.e.m., while insets are distribution box plots of the same parameter. In (B) numbers in the bars are # of fields of view analyzed/# of animals analyzed, while for (C-I) they are # of trajectories/# of animals analyzed. (B) Fraction of moving particles, with directed motion, showing no difference between WT and MUT ($p = 0.83$). (C) Directionality. Left : pie charts of the percentage of purely anterograde (A), purely retrograde (R), or bidirectional trajectories (B), for each condition. The proportion of purely A trajectories decreases in favor of purely R trajectories in MUT. Right : fraction of R transport in bidirectional trajectories. MUT shows a bias towards retrograde transport ($p = 0.018$). (D-E) Retrograde and anterograde segmental velocities do not vary between conditions ($p = 0.896$ and $p = 0.181$, for R and A respectively). Trajectories with purely anterograde or purely retrograde movement were excluded from analysis of retrograde or anterograde segmental velocities, respectively. (F) Pausing frequency, that does not vary ($p = 0.574$). (G) Pausing duration decreases by 16% ($p = 0.046$). (H) The average retrograde run length is 68% longer for MUT compared to WT ($p = 0.00092$). Trajectories with purely anterograde movement were excluded from analysis. (I) Anterograde run length does not vary ($p = 0.303$). Trajectories with purely retrograde movement were excluded from analysis.

to 14.2%, the fraction of pure A trajectories is only 3.4% in the MUT to the benefit of pure R trajectories which increases from 7.5% to 10.8%. As for the study of DYNA effect, we also focused on the remaining directionality in the largest fraction of bidirectional trajectories. We evaluated the fraction of retrograde transport in this sub-population and found that it is broadly distributed (inset of Figure 5C) indicating that bidirectional trajectories are largely polarized. Moreover, this fraction also displays a median shift from 0.51 to 0.61 between WT and MUT, which means that bidirectional trajectories are biased towards the retrograde direction in mutants.

Furthermore, as observed in the case of dynein inhibition with DYNA, in *kif5aa* mutant we did not detect significant changes of both A and R segmental velocities that remain of 0.8-0.9 $\mu\text{m/s}$ (Figure 5D-E). The pausing frequency does not vary statistically neither between WT and MUT, but we observe a decrease of the pausing duration (all types of pauses gathered) of about 15% from 0.382 to 0.323 s. Decreases were also observed for all pause subcategories (*i.e.* between two identical directions or two opposite directions of motion), but none of them was individually significant (Fig.S9D-F). Moreover, we observe a significant increase of the retrograde RL by 68% from 347 to 585 nm between WT and MUT, without effect on the anterograde RL (Figure 5H-I).

Altogether, the directionality study and endolysosomal transport parameters in *kif5aa*

larvae indicate that reducing Kif5aa concentration biases the direction of transport towards the retrograde direction and favor longer R run lengths. These results are compatible with the tug-of-war model of intraneuronal transport,²³ stating that (i) both types of R and A motors (dynein and kinesins respectively) are present on the cargo, (ii) that they are both attached to the microtubule pulling in opposite directions, and (iii) that the direction of actual motion is dictated by the set of active bound motors pulling the strongest, which is likely to be the dynein motors in our situation of reduced amount of Kif5aa kinesin 1 motor.

The absence of modification of anterograde velocity (Figure 5E) and more surprisingly of anterograde run length (Figure 5I) in *kif5aa* mutants is in favor of a compensation mechanism. In Zf larvae such compensation can occur by other Kif5 motors to ensure anterograde motion, or by another kinesin motor complex of the superfamily, such as kinesin 2 motor complex including Kif3a/3b/Kap3 and Kif3a/3c/Kap3,⁴⁹ which is known to drive vesicle axonal transport too.^{23,50} In the case of inhibitor experiments, as dynein heavy chain 1 and 2 isoforms are the only motors ensuring retrograde transport, and as they were reported to be both inhibited by dynapyrazole A *in vitro*,³³ no motor substitution is expected to compensate their inactivation. However, at the dynapyrazole concentration used, the retrograde transport is not fully halted, and the overall transport displays only a reduced retrograde run length (Figure 4H) and increase in pausing frequency (Figure 4F), with no modification of velocity (Figure 4E). These observations may reveal a limited intravital penetration of dynapyrazole A, but we could not test this hypothesis because of the toxicity observed for higher concentration (possibly due to dynein 2 concomitant inhibition).

NanoKTP polar angle fluctuations measurement. Finally, we took advantage of the blinking of the SHG signal from nanoKTP to reveal modifications in the endosome/lysosome-motor-microtubule binding in *kif5aa* mutants. Indeed, this blinking reflects changes of orientation of the vesicle relative to the microscope objective axis (Figure 1C), as the nanocrystal is presumed immobile relative to its embedding vesicle.²⁹ To quantify this fluctuation, we first reverted from $P_{2\omega}^{\text{det}}(\theta)$ (the detected SHG power at frequency 2ω), the polar angle θ between the direction of light propagation and nanoKTP *c*-axis (which is also the direction of the equivalent SHG-emitting dipole, see Figure 1C). To this aim we calculated the efficiency $\eta(\theta)$ of the SHG power signal collection through the microscope objective in function of θ , and obtained the general form $\eta(\theta) = 3/8\pi(B \sin^2 \theta + C)$, where B and C are parameters depending only on the microscope objective NA (Supporting Information Data S10). SHG detected power differs from SHG collected one $P_{2\omega}$ by a constant coefficient taking into account optical losses and the detector quantum efficiency. Figure 2A shows that all SHG signals from nanoKTP-labeled vesicles with directed motion vary between almost zero power $P_{2\omega}^{\text{det},\text{min}}$ and a maximum $P_{2\omega}^{\text{det},\text{max}}$ that depends on the particle size. We assumed that these extreme power values correspond to $\theta = 0^\circ$ and 90° respectively, and then inferred θ at each trajectory point from mapping the detected power to $\eta(\theta)$ by $P_{2\omega}^{\text{det}}(\theta) = (P_{2\omega}^{\text{det},\text{max}} - P_{2\omega}^{\text{det},\text{min}}) \sin^2 \theta + P_{2\omega}^{\text{det},\text{min}}$. Note that we are not able to extract the azimuthal angle defining the orientation of the SHG dipole in the sample plane, as we are not resolving the polarization of the collected light.

Then in order to quantify the fluctuations of vesicle orientation, we first applied a Savitzky-Golay filter to SHG intensity time traces in order to mitigate the high sensitivity of the intensity-to-polar angle inversion process to a large intensity fluctuation close to

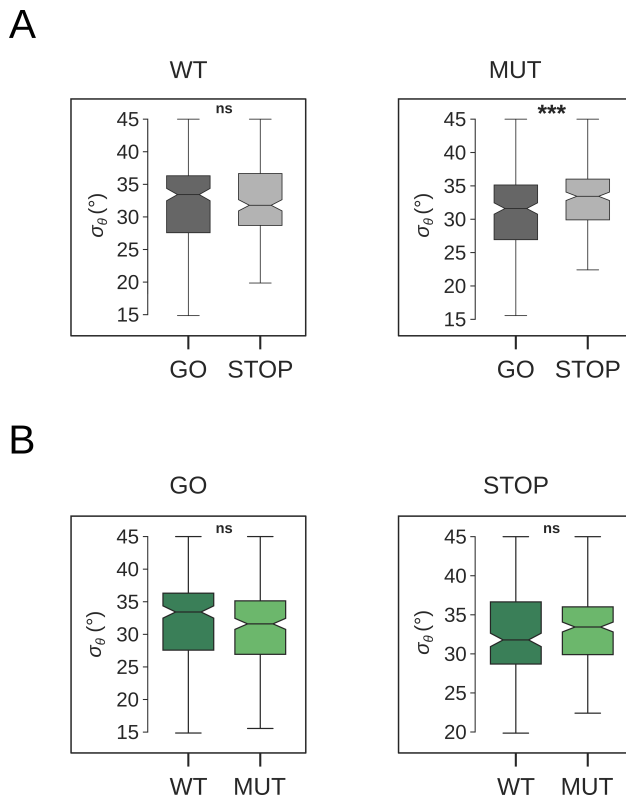


Figure 6: Fluctuation of the orientation of nanoKTP-labeled vesicles during Go and Stop phases of motion in neurons of the brain of *kif5aa* Zf larvae, as quantified with the standard deviation σ_θ of the polar angle θ . (A) Comparison of the fluctuations between the Go and Stop phase for the two conditions, showing that σ_θ has no difference between Go and Stop phases ($p = 0.99$), but the fluctuations are larger in the Stop (average of 33.1°) compared to the Go (30.6°) in the mutant ($p = 1.5 \times 10^{-4}$). (B) Comparison of σ_θ between the two conditions in the different phases, showing no difference between WT and MUT in the Go ($p = 0.06$) and Stop ($p = 0.07$) phases.

the extrema. We determined $\theta(t)$ from the filtered intensity time trace and calculated its standard deviation σ_θ in each phase of motion within a trajectory, from which we inferred an average value $\overline{\sigma_\theta^{\text{Go/Stop}}}_j$ per trajectory j . We focused our analysis on *kif5aa* data set. Figure 6A displays the distribution of these values for all the trajectories of WT and MUT conditions. We notice that in the MUT case $\{\sigma_\theta^{\text{Stop}}\}$ distribution (simply noted $\sigma_\theta^{\text{Stop}}$ after) is significantly larger by 8% than the one of $\sigma_\theta^{\text{Go}}$ and that there is no difference in the WT case. Comparing now $\sigma_\theta^{\text{Go}}$ and $\sigma_\theta^{\text{Stop}}$ between WT and MUT, we did not observe differences of orientational fluctuations (Figure 6B). To our knowledge, such fluctuations of endolysosomal vesicle orientation were not reported *in vivo* before. In the case of the mutant, our data are in agreement with observations performed *in vitro* in cultured PC12 rat neuronal cell line showing larger orientational fluctuations of the vesicle in Stop than Go phases, as detected from differential interference contrast signal from embedded gold nanorods.²⁸ However, in primary cultures of dorsal root ganglion rat neurons, Kaplan et al.,²⁹ did not observe any differences of the rotational lability (another measure of the rotational fluctuations) between

the Stop phases and Go phases, like in the case of *kif5aa* WT siblings.

In order to check the consistency of our observations in different conditions, we also computed the vesicle orientation fluctuations for the dynapyrazole dataset (Supporting Figure S12). We observed similar distributions and median values of σ_θ than in *kif5aa* data, with larger $\sigma_\theta^{\text{Stop}}$ than $\sigma_\theta^{\text{Go}}$ for both control and DYNA. Here, the inhibition of functional dyneins by dynapyrazole leads to an increase of 14% of σ_θ in the Go phase, and no changes in the Stop ones (Figure S12B).

Conclusions

In this work we described a method for *in vivo* quantification of axonal transport in neurons of the CNS, based on the measurement of a large set of parameters that are able to reveal transport impairments finely. This method relies on intravital fast two-photon microscopy recording of the motion of axonal endolysosomal vesicles, thanks to their labeling with non-linear nanocrystals of KTP, as a first step, followed by a high-throughput data processing extraction of endosome trajectories and associated transport metrics. We established this approach *in vivo* by using the zebrafish model organism, exploiting its translucency and genome editing possibilities. We achieved sparse labeling of endolysosomal vesicles traced in the axonal compartments of CNS neurons through microinjection of the nanoKTP in the optic tectum of Zf larvae. About 90% of the injected larvae had, in the region of interest, a concentration of nanoKTP well suited to isolate single trajectories after the nanocrystals spontaneous axonal uptake.

This easy to apply original approach contrasts with the use of transient knock-in consisting in the injection of nucleic acid sequences encoding fluorescent proteins and sorting of the larvae presenting the relevant phenotype. Moreover, the nanoKTP are well tolerated by the Zf larvae and exhibit a large SHG signal that is perfectly photostable, in contrast to conventional molecule-based fluorophores that bleach during the recording. Our approach allowed us to reach an average resolution of 129 nm at 20 frames per second, which is a photon budget-limited compromise between high spatial resolution and fast acquisition. This spatiotemporal resolution achieved in our method has no equivalent in *in vivo* studies of axonal transport measurements to date.⁶ Therefore it constitutes a major advance to precisely quantify slow (Stop) phases-related parameters and to detect altered cargo pausing associated to abnormal neuronal activity. In addition, the intracerebral injection in the optic tectum of the Zf brain led to an efficient uptake of nanoKTP localized in the axons of the peri-ventricular neurons, allowing to study independently kinesin or dynein driven directions of motion.

To demonstrate the sensitivity of our *in vivo* assay, we probed the impact of reducing the concentration of the active molecular motors on the transport dynamics. To this aim we reduced the concentration of dynein (using dynapyrazole, the best-in-class dynein inhibitor) or kinesin 1 Kif5aa (using *kif5aa* loss-of-function zebrafish mutant line) which is a member of kinesin 5Aa family, abundantly expressed in neurons. We observed differential overall effects of these modifications: dynapyrazole reduces by 32% the mobile fraction of nanoKTP directed motion, proving its efficiency as a dynein inhibitor in Zf larvae brain. No difference in this fraction was detected between wild type and mutant *kif5aa* larvae.

Dynapyrazole also consistently reduces the retrograde run length by 37%. *Kif5aa* mutants showed a 68% increase of the retrograde run length, instead of a reduction of the anterograde run length. To interpret this observation, we hypothesize that other kinesins might replace Kif5aa and maintain the anterograde transport. As such, kinesin 2 is one possible candidate as it binds more loosely to the microtubule,⁵¹ which could explain the larger orientational fluctuations of the endolysosomal vesicle-containing nanoKTP in mutant *kif5aa*. However, further experiments are required to validate this interpretation.

In our current assay, the spontaneous endocytosis of bare nanoKTP in axonal compartments could be enhanced by conjugating transferrin or rabies virus glycoprotein peptides to nanoKTP. Future developments will also extend the use of our method to other neuronal cell subtypes and/or later stages of development. Taking advantage of the low absorption of tissues in near infrared excitation wavelength, and the possibility to easily tune excitation wavelength minimize the tissue auto-fluorescence contribution, our method should allow to image more complex and thicker biological model, such as juveniles or adults zebrafish mutants devoid of pigmented cells.⁵² These approaches will provide a way to study axonal transport in physiological conditions at different stages of development in normal and pathological situations. The latter include influence of intrinsic (*i.e.* mutations inducing transport defects) or extrinsic (*i.e.* temperature, water quality, chemical exposure. . .) factors.

Finally, our assay could also be valuable to develop biomedical models reproducing human nervous system alterations quantitatively assess the link between axonal transport impairment and disease outcome. In particular, it would be interesting to reproduce our experiments in Zf larvae infected by neurotropic viruses,⁵³⁻⁵⁵ in transgenic lines used as models of neurodegenerative diseases,⁵⁶⁻⁵⁸ or upon axonal photoablation⁵⁹ to precisely measure the axonal transport impairments in these situations. In case of detection of a significant impairment of axonal transport, our assay could be harnessed to screen therapeutic compounds able to rescue the defect.

Materials and Methods

Zebrafish lines and husbandry

Wild-type (AB strain) zebrafish (*Danio rerio*) were raised in the IERP fish facility at INRAE Jouy-en-Josas, France. *kif5aa*^{*162} was generated as previous described.³² The *dync1h1*^{mw20} mutant line was described by Insinna *et al*³¹ Zf were maintained at 28°C on a 14 h light/10 h dark cycle. After spawning, eggs were collected and incubated at 28°C in fish system water supplemented with 0.3 µg/mL methylene blue. PhenyIthiourea (PTU) was added at 24 h postfertilization to inhibit synthesis of melanin that would absorb the imaging excitation laser light and harm the animal.

Ethics statement Zebrafishes were housed in the animal facility of our laboratories which were built according to the respective local animal welfare standards. All animals were handled in strict accordance with good animal practice as defined by the European Union guidelines for the handling of laboratory animals (http://ec.europa.eu/environment/chemicals/lab_animals/home_en.htm). All animal procedures were performed in accordance with French and European Union animal welfare guidelines with protocols approved by committees on ethics of animal experimentation of Sorbonne Université (APAFIS#21323-2019062416186982) and Université Paris-Saclay. All animal work was also approved by the Direction of the Veterinary Services of Versailles, France (authorization number C78-720), and experimental protocols were approved by the INRAE institutional ethical committee “Comethea” (APAFIS#22160-2019092615256610).

Genotyping of *dync1h1*^{mw20} and of the *kif5aa*^{*162} mutant alleles

For *kif5aa*^{*162} genotyping the following primers were used (5' to 3'): *kif5aa* geno fwd: GTTCACAGATTGTGATGTCTGTG, *kif5aa* geno rev: TGGAGGATG- GAGAAATGATGACA. After PCR amplification from genomic DNA the 400 bp long amplicon was digested with NcoI. The wild-type allele is digested into two fragments of 240 bp and 160 bp length, respectively. The mutant homozygote alleles is not digested and show a single band at 387 bp or 390 bp, while the heterozygote genotype is characterized by three bands at 160, 240 and 390 bp. For *dync1h1*^{mw20} genotyping the following primers were used (5' to 3'): *mw20* geno fwd: CACGAGGAGCTCTACAAGTGG, *mw20* geno rev: GAACAGGTTG-GCGTAGTGGT. After PCR amplification from genomic DNA the 750 bp long amplicon was digested with RsaI. The wild-type allele is digested into two fragments of 350 bp. The mutant homozygote alleles is not digested and show a single band at 750 bp, while the heterozygote genotype is characterized by two bands at 750 and 350 bp.

KTiOPO₄ nanocrystal synthesis and size separation KTP nanoparticles were synthesized following a protocol adapted from Ref.²⁶ In a typical experiment, a titanium alkoxide solution acidified with HCl is first mixed with a solution of monobasic potassium phosphate leading to the formation of a precipitate. The solution is then neutralized up to pH=6 through the addition of K₂CO₃. The precipitate is then recovered by centrifugation, washed several times with water and dried. The powder is then thermally treated to ensure KTP phase formation with an optimal temperature of 700° for 2 hours. The powder is then washed

three times by centrifugation with deionized water and the pellet redispersed in water leading to the colloidal suspension of KTP nanocrystals.

The hydrodynamic particle size for this pristine suspension was estimated to be 230 nm by dynamic light scattering (DLS, volumic fraction). As previous characterization have shown a significant size dispersion,²⁶ we decided to apply a size selection procedure to narrow the size distribution. To this aim, the pristine suspension was first centrifugated at an acceleration of 8,000 *g* for 5 min. Then, the supernatant was extracted and centrifuged at 60,000 *g* for 5 min. The supernatant was discarded and the pellet was redispersed in pure water leading to the colloidal suspension used in this work. DLS analysis of this solution gave a cumulant *Z*-average particle size of 126 nm (volumic fraction analysis), confirmed by nanoparticle tracking analysis (Supporting Data S1) yielding a value of 123.8 ± 2.4 nm.

Microinjection of nanoparticles in zebrafish larvae Before injection, larvae were anaesthetized with eugenol diluted at 0.0075% in fish system water supplemented with methylene blue and PTU as described above. Larvae were injected in the left brain hemisphere with 5 nl of nKTP solution using pulled borosilicate glass microcapillary (GC100F-15, Harvard Apparatus) pipettes under a stereomicroscope (MZ10F, Leica Microsystems) with a mechanical micromanipulator (M-152; Narishige), and a Femtojet microinjector (Eppendorf).

To investigate the transport properties of lysosomes and the colocalization of moving nanoKTP with these compartments, 4 dpf zebrafish larvae were exposed during 1 hour to LysoTracker Red DND-99 (Ref.L7528, Invitrogen/ThermoFisher) diluted at 5 μ M in fish system water supplemented with methylene blue and PTU. Larvae were rinsed briefly before mounting for live imaging.

Immunostaining Zebrafish larvae were fixed overnight at 4°C in PBS 0.01 M + 4% formaldehyde. Larvae were rinsed three times for 5 minutes with PBS + 0.1% tween. All the following steps were performed at room temperature (RT) unless indicated otherwise. Larvae were dehydrated in ascending methanol series (20%, 40%, 60%, 80% in H₂O) during 15 minutes for each step. Larvae were incubated in 100% methanol overnight at 4°C. Dehydrated larvae were rehydrated in descending methanol series (80%, 60%, 40%, 20% in water) during 15 minutes for each step. Samples were permeabilized by immersion in PBS + 0.5 μ M CaCl₂ + 1 mg/ml collagenase (C9891, Sigma-Aldrich) during 15 minutes. Larvae were rinsed three times during 5 minutes in PBS + 0.2% triton X100. Larvae were then blocked in PBS + 5% triton X100 + 10% DMSO + 10% horse serum + 0.05% sodium azide for 5 hours at 37°C. Mouse anti-acetylated tubulin (T7451, Sigma-Aldrich) and rabbit anti-GFAP (ZO334, Dako) were both diluted at 1/500 in PBS + 5% Triton + 10% DMSO + 10% horse serum + 0.05% sodium azide and applied 2 days at 37°C. After repeated washes in PBS + 5% triton + 10% DMSO + 0.05% sodium azide, samples were incubated overnight at 37°C with Alexa Fluor 488 goat anti-mouse (A11001, ThermoFisher) and Alexa Fluor 594 goat anti-rabbit (A11012, ThermoFisher,) diluted at 1:500 in the same solution than first antibody was applied. Larvae were rinsed several times with PBS + 0.1% tween. Before imaging, larvae were cleared by incubation in RIMS overnight.⁶⁰ Larvae were mounted under #1 coverslips in RIMS supplemented with 0.8% low-gelling agarose.

nonlinear microscopy live imaging and immunostaining imaging nonlinear live imaging of zebrafish larvae was performed with an upright Leica SP8 two-photon microscope using a HCX IRAPO 25X/0.95NA water immersion objective (Leica Microsystems). Anaesthetized larvae were transferred to 3% agarose casts containing wells allowing to position larvae. Larvae were then embedded in 0.5% low-gelling agarose. Orientation of larvae was adjusted to dorsal view before complete gelation. Fish system water + methylene blue + PTU + eugenol was added on embedded larvae. NanoKTP were excited at 1040 nm with a Chameleon Vision II laser (Coherent). A quarter-waveplate was used to convert the linear excitation polarization into a circular one. The emitted photons were detected in a non-descanned (NDD) pathway with a Gallium Arsenide Phosphorous-based hybrid detector (HyD, Leica), having 50% quantum efficiency at 520 nm wavelength, and a maximum counting rate of 60 Mcounts/s in the conventional photon counting mode, extended to 300 Mcounts/s in the “standard” mode that we used, and that compensate for the nonlinearity of conventional counting. We put a 525/50 nm bandpass (green channel) in front of the HyD. Temperature was maintained constant during and between experiments by using an incubator box combined with a precision air heater (The Cube, Life Imaging Service). All experiments were performed at 24°. For the live imaging of LysoTracker Red with two-photon excitation, we tuned the pulsed laser wavelength at 1030 nm, as 1040 nm corresponds to a minimum of the two-photon absorption of this dye.⁶¹ This had no impact on nanoKTP SHG detection as its spectrum, now centered on 515 nm with 10 nm FWHM, still falls within the 525/50 nm bandpass filter of the green channel. We detected the LysoTracker Red emission (red channel) on a similar HyD as the green, placed on the transmission path of the beamsplitter (edge at 560 nm wavelength) separating green and red channel, and preceded by a 585/40 nm bandpass filter.

Immunostained samples were acquired by confocal and 2-photon microscopy using the same Leica SP8 and a coverslip corrected HCX IRAPO L 25X/0.95NA water immersion objective (Leica Microsystems). Alexa Fluor 488 and Alexa Fluor 594 were excited with lasers emitting at 488 nm and 552 nm wavelengths respectively. Fluorescence was detected with PMT detectors. SHG signal from nanoKTP in fixed larvae was acquired as in live imaging experiments.

Automatic extraction of nanocrystal-labeled vesicle trajectories and transport parameters Videos were first exported from the Leica LIF proprietary format as stacks of 8-bits TIFF files with a custom Fiji macro relying on the import function of the Bio-Formats plugin.⁶² The subsequent data analysis process is automated, using a set of Python scripts that we developed and made freely accessible online.³⁶ It consists of two main parts: Part 1 aims at extracting valid trajectories directly from the video files and Part 2 calculates transport parameter observables from the trajectories.

Part 1 is composed of three steps: 1) raw trajectories extraction from the videos, 2) filtering directed motions only, 3) regularisation of the localization noise. The stacks are first preprocessed by a white top-hat transform, then processed by the `trackpy`⁶³ implementation of the Crocker-Grier algorithm.⁶⁴ This algorithm locates features of a given size (in pixels), brightness (in counts) and minimum separation (in pixels) in each frame (we selected the following parameters of the `batch` function: `diameter=9`, `minmass=300` and `separation=12`),

then links them to form $(x(t), y(t))$ trajectories, as paired arrays of coordinates over time t . To this aim we used the `link` function with the following parameters: `search range=6`, `memory=5`, `adaptive stop=5` and `adaptive step=0.9`. These trajectories are then filtered according to their shape, based on a Mean Square Displacement (MSD) threshold of 300. Trajectories with MSD lower than this threshold were discarded. This filters out nonlinear, aberrant trajectories. We found that this algorithm sometimes resulted in a visually singular trajectory being broken down into several smaller ones. We thus rejoined trajectories of which beginnings and ends are within a given spatiotemporal range from one another. The rejoining thresholds were 10 frames and 40 pixels, *i.e.* the end of one trajectory had to be within 10 frames and 40 pixels distance of the beginning of another one for them to be rejoined. As we aim to measure directed axonal transport, ideal trajectories should progress in a relatively rectilinear motion. Hence, we first rotated each trajectory $(x(t), y(t))$ in a way that the line between the first and furthest points becomes horizontal, and then only selected trajectories which does not deviate from their fit to a third degree polynomial function $f(x)$ more than 1.4 pixels in standard deviation, *i.e.* trajectories for which $\sqrt{\langle (y - f(x))^2 \rangle} < 1.4$ pixels, where $\langle \dots \rangle$ is the average on all trajectory points.

The set of selected trajectories was then subjected to a noise filtering, based on an algorithm of convex minimization of acceleration, in order to compensate for localization uncertainty. A measured raw trajectory $\mathbf{r}_m(t)$ extracted from the videos by `trackpy` consists of the underlying directed motion $\mathbf{r}_d(t)$ of vesicles moved by the molecular motors, onto which a measurement noise is superimposed that we modelize as a white Gaussian noise $\mathbf{w}_G(t)$:

$$\mathbf{r}_m(t) = \mathbf{r}_d(t) + \mathbf{w}_G(t).$$

In order to extract the ground truth vesicular motion $\mathbf{r}_d(t)$ we must correct for the noise first. The vesicular displacements being composed of a succession of phases of almost constant velocity, interrupted by rare pauses with almost zero velocity, the correction implemented relies on the discrimination of these rare but consequent changes in velocity from the permanent ones caused by the noise. An efficient approach to perform this denoising in such a situation is to apply a total variation reconstruction (TVR).⁴¹ Briefly, we are looking for large changes in the value of the measured (noisy) velocity. Considering $\hat{\mathbf{r}}$ as an estimator of $\mathbf{r}_d(t)$, our TVR approach consists in minimizing the acceleration $\|D^2\hat{\mathbf{r}}\|_1$ where D is the derivation operator relative to the time and 1 stands for the $l1$ -norm, under some constraints. The latter is that the estimator $\hat{\mathbf{r}}$ must remain close to the acquired trajectory $\mathbf{r}_m(t)$. With our hypothesis on the noise, we can write that $\|\mathbf{r}_m - \mathbf{r}_d\|_2^2 \approx n\bar{\sigma}_{xy}^2$, where $\bar{\sigma}_{xy}$ is the average standard deviation of the Gaussian noise $\mathbf{w}_G(t)$, n is the number of frames in the trajectory, and 2 stands for the $l2$ -norm.

This leads to the following convex minimization problem:

$$\begin{aligned} & \text{minimize } \|D^2\hat{\mathbf{r}}\|_1 \\ & \text{subject to } \|\hat{\mathbf{r}} - \mathbf{r}_d\|_2^2 \leq n\bar{\sigma}_{xy}^2 \end{aligned}$$

To solve this problem, we use `cvxpy` (version 1.1.7)^{65,66} Python package that relies on the convex optimization `SCS` solver^{67,68} (version 2.1.2).

Part 2 consists in parsing the trajectory into Go and Stop phases, corresponding to fast and little to no movement, respectively, using the same confinement ratio (R) method

used in our previous work.²² We determined the threshold $R_{\text{th}} = 0.64$ from the minimum of its distribution when considering all trajectories (Supporting Figure S5). At each point i we calculate R_i (from 3 preceding and 3 successive positions): $R_i > R_{\text{th}}$ is associated to a Go phase, while $R_i \leq R_{\text{th}}$ is a Stop phase. As the field of view encompasses a very polarized region, with axons projecting into the neuropil, we can discriminate anterograde and retrograde transport based on which direction the particle is moving. Several parameters are then calculated, including the general parameter of the *fraction of moving particles* and five trajectory-specific parameters:

- **Fraction of moving particles:** ratio of moving particles to non-moving ones. It is calculated by dividing the number of trajectories analyzed for each file (*i.e.* one field of view) by the number of particles found on the first frame of that file. It does not take into account trajectories that were filtered out before analysis, or particles that might appear after the first frame. It is therefore not an absolute measure of the fraction of moving particles, and is only used for relative comparison between experimental conditions.
- **Directionality:** sum of the lengths of the retrograde segments divided by the sum of the absolute values of the lengths of all segments (anterograde and retrograde). A fraction value of 1 correspond to a purely retrograde trajectory, and 0 to a purely anterograde one.
- **Segmental velocity:** speed of the endolysosomal vesicle-embedding particle in $\mu\text{m}/\text{s}$. It is calculated as the average speed of Go phases, which is itself the average of the point velocities for each phase. It is evaluated separately for retrograde and anterograde directions.
- **Pausing frequency:** frequency at which the particle pauses, in number of events per minute. It is calculated as the number of stops divided by the total trajectory time.
- **Pausing duration:** Average duration, in seconds, that the particle spends in Stop phases. It is calculated as the average duration of Stop phases in a given trajectory.
- **Run length:** average length, in μm , traveled during Go phases. It is calculated as the average length of Go phases of a trajectory, which is itself the segmental velocity of each phase multiplied by its duration.
- **Theta angle fluctuations σ_θ :** standard deviation, in $^\circ$ of the polar angle θ calculated in Go or Stop phases. The angle is first calculated point by point from the intensity (Supporting Data S8, Figure S10), then the standard deviation of that angle for each phase is averaged per trajectory.

Statistical analysis and data representation Statistical differences were analyzed by Wilcoxon signed-rank test (except in Fig. 3, see associated caption). * corresponds to $p < 0.05$, ** $p < 0.01$ and *** $p < 0.001$. All bar plots represent the average of each parameter value, with error bars being \pm standard error of the mean (s.e.m.). In all box plots the box delimits the first and third quartiles and the whiskers extend from the first to the ninth

decile (10-90%). Outliers are not shown. The median line is surrounded by a notch of which the width defines a confidence interval.

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Associated content

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Supporting Information Available

The following files are available free of charge:

- Supporting information file containing data and figures on: nanoKTP electron microscopy images and size distribution; emitted light spectrum; method of conversion

of 8 bits-encoded hybrid detector intensity levels in photon counts; precision of localization of the nanoKTP; determination of confinement ratio threshold to parse the trajectory in Go and Stop phases; proofs of the absence of effect of nanoKTP on lysosomes axonal transport; analysis of colocalization of nanoKTP with lysosomes; reversal of directionality quantification for dynapyrazole-treated larvae and *kif5aa* larvae; separate quantification of pausing duration for pauses between two anterograde or two retrograde phase of motion, or between phases of opposite directions; collection efficiency of SHG light emitted by a single monocrystalline nanoKTP; example of $\theta(t)$ retrieval from SHG intensity variation, including the case of static nanoKTP; fluctuations of orientations of nanoKTP-labeled endosome endolysosomal vesicle in dynapyrazole-treated zebrafish larvae.

- SupportingVideoS1.avi: raster-scanning video from which Fig. 2A was calculated, showing in inverted grey scale the SHG signal of nanoKTP moving in PVN neuron of a zebrafish larva (*kif5aa* wild-type). True frame rate of 20 frames/s, total of 2354 frames. Scale bar: 10 μm .
- SupportingVideoS2.avi: another raster-scanning video showing in inverted grey scale the SHG signal of nanoKTP moving in PVN neuron of a hetegozygous *kif5aa* zebrafish larva. True frame rate of 20 frames/s, total of 2501 frames. Scale bar: 10 μm .
- SupportingVideoS3.avi: example of a raster-scanning video of nanoKTP immobile on a glass coverslip, acquired in the same conditions as the zebrafish larvae, and used to estimate the precision of localization (Figure S4C).

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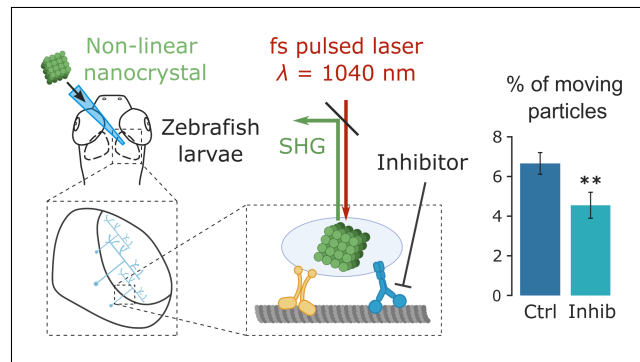
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TOC Graphic



We measured transport parameters of endolysosomal compartments in axons of zebrafish larvae brain. To this aim we injected nonlinear nanocrystals (NC) and traced their motion after their spontaneous endocytosis, by recording their second harmonic generation (SHG) excited with near infrared femtosecond (fs) laser pulses. The quantitative analysis of NC-labeled endolysosomes motion is sensitive enough to reveal the inhibition of molecular motors transporting these compartments.