

Optical control of fast and processive engineered myosins in vitro and in living cells

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Precision tools for spatiotemporal control of cytoskeletal motor function are needed to dissect fundamental biological processes ranging from intracellular transport to cell migration and division. Direct optical control of motor speed and direction is one promising approach, but it remains a challenge to engineer controllable motors with desirable properties such as the speed and processivity required for transport applications in living cells. Here, we develop engineered myosin motors that combine large optical modulation depths with high velocities, and create processive myosin motors with optically controllable directionality. We characterize the performance of the motors using in vitro motility assays, single-molecule tracking and live-cell imaging. Bidirectional processive motors move efficiently toward the tips of cellular protrusions in the presence of blue light, and can transport molecular cargo in cells. Robust gearshifting myosins will further enable programmable transport in contexts ranging from in vitro active matter reconstitutions to microfabricated systems that harness molecular propulsion.

ngineered cytoskeletal motors with dramatically altered properties and controllable functions¹ have provided critical tests of motor mechanisms^{2,3} and tools for relating nanoscale biophysics to cellular and tissue-scale functions^{4,5}. An outstanding challenge is that engineered motors often have impaired parameters, such as speed and processivity⁶⁻⁸, in comparison with their natural counterparts. Eukaryotic cells rely on both processive and non-processive cytoskeletal motors with speeds ranging from the nanometer per second to micrometer per second range9. Light-activated gearshifting10 provides direct control over cytoskeletal motor speed and direction in genetically encoded engineered proteins, and has been proposed to be applicable for regulating cellular dynamics. This technology has been demonstrated in vitro, using gliding filament assays with purified proteins. However, existing designs for light-activated gearshifting¹⁰ are non-processive and suffer from either low overall speeds or modest velocity modulation, precluding their use in many areas of cell biology, devices and reconstituted systems. Moreover, unlike complementary optogenetic approaches based on cargo recruitment¹¹⁻¹⁴, gearshifting has not been shown to function in cellular contexts.

In this Article we demonstrate that engineered myosin motors can satisfy four critical design constraints for optically controllable transport: speeds that match the range of speeds of natural cytoskeletal motors, deep modulation in response to light, processive motility and operability in living cells. Exploiting the principle of programmable changes in lever arm geometry 67,10,15, we first introduce fast non-processive myosin motors that respond strongly to blue light in gliding filament assays. Collections of these motors show graded dose-dependent responses to optical stimulation. We then use

in vitro single-molecule fluorescence imaging and gold nanoparticle tracking to characterize the motility of tetrameric engineered motor assemblies, and we find that these designed motors walk processively along actin filaments and switch their predominant direction of stepping in response to blue light. Moving to live-cell imaging, we observe light-dependent intracellular localization of the bidirectional tetramers in both fibroblasts and cultured neurons, and confirm high-speed motility of individual motors tagged with fluorescent protein arrays. Finally, we use co-expression in epithelial cells to show that the engineered myosins can be used to transport separate molecular cargos, demonstrating light-dependent localization of both cytoplasmic and transmembrane proteins to cellular protrusions.

Results

Engineering fast myosins with strong optical response. To enable cellular applications that require robust spatiotemporal control while matching or exceeding the transport velocities of natural motors, we first set out to design a gearshifting myosin that combines high velocity with deep optical modulation. The existing MyLOV family of engineered gearshifting myosins ¹⁰ includes constructs (MyLOV1–4) that show deep light-dependent velocity modulation but low speeds ($<10\,\mathrm{nm\,s^{-1}}$), and one construct (MyLOVChar) with high speed ($\mu\mathrm{m\,s^{-1}}$) but only mild velocity modulation. We hypothesized that a suitable starting point for optimization could be MyLOVChar, a light-controlled motor incorporating the catalytic domain of *Chara corallina* myosin XI, the fastest known cytoskeletal motor ¹⁶.

To identify possible strategies for achieving our design goals, we reviewed the switching mechanism and functional characteristics

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of MyLOVChar. The MyLOVChar design exploits a mechanical effect of light-induced unfolding in the N-terminal Jα-helix of a LOV domain (Avena sativa LOV2)17 embedded in an engineered lever arm structure (Fig. 1a). The fully folded LOV domain is designed to act as a mechanical hairpin element that redirects the lever arm in the absence of blue light, resulting in a small stroke of the myosin towards the actin (+)-end. In the light, the J α -helix is undocked from the LOV domain and becomes disordered. The last rigid element of the LOV domain then becomes the effective end of the lever arm, predicted to produce a larger (+)-end directed stroke and an increased resulting velocity^{18,19}. A coarse structural model of the MyLOVChar power stroke estimated that the velocity should increase by more than threefold in the light¹⁰. We used cryo-EM (Extended Data Fig. 1a-c and Supplementary Table 1) to obtain a new experimental post-stroke structure of Chara corallina myosin XI bound to actin (Fig. 1b, Extended Data Fig. 1d and Supplementary Video 1) and generated updated models of MyLOVChar, producing a similar prediction with even larger velocity modulation (Extended Data Fig. 2a). However, as reported previously, the experimentally measured modulation is much smaller. In an in vitro gliding motility assay (Fig. 1c), the speeds of actin filaments driven by MyLOVChar increased from ~3-5 µm s⁻¹ in the absence of blue light to $\sim 5-7 \, \mu m \, s^{-1}$ when illuminated (Fig. 1d and Supplementary Video 2), for an optical modulation depth (defined as $1 - v_{\rm dark}/v_{\rm lit}$, with $v_{\rm dark}$ and $v_{\rm lit}$ determined as shown in Fig. 1e) of ~35%. This discrepancy between prediction and experiment may be explained by unintended compliance or angular deviations introduced by junctions between structural modules in the engineered lever arm, leading to a power stroke that is suboptimally redirected in the dark. We pursued two complementary strategies for overcoming this deficiency: compensating for poor redirection by increasing the length of the redirected lever arm, and correcting suboptimal junctions by varying the junction sequences as in previous work¹⁰.

In the first strategy, we aimed to compensate for compliance in the MyLOVChar lever arm by adding spectrin repeats as rigid extensions that increase the length of the redirecting part of the lever arm. MyLOVChar2 incorporates four added spectrin repeats (Supplementary Fig. 2) and displays a reduced gliding velocity in the dark, as expected for a more redirected lever arm (Fig. 1d and Supplementary Video 3). The lit velocity is also reduced relative to MyLOVChar, but to a smaller degree, so that the net optical modulation is increased to ~60% (Fig. 1f). Thus, building longer redirecting elements is one effective strategy to compensate for mechanical inefficiency in gearshifting myosins.

In the second strategy, we aimed to improve the mechanical efficiency of the design by optimizing the mechanical properties of the junction between the LOV domain and the flanking spectrin repeats. The resulting construct MyLOVChar3 (Fig. 1d and Supplementary Video 4) uses a junctional design adapted from MyLOV1, the junctional variant with the strongest redirected stroke from the MyLOV1–4 series of slow gearshifting motors based on porcine myosin VI¹0. MyLOVChar3 achieves an increased optical modulation of ~50%, showing that optimizing the junctions of engineered domain fusions can also improve the mechanical efficiency of gearshifting lever arm designs.

Next, to maximize the optical modulation of the velocity, we combined the two successful strategies, retaining the improved junction from MyLOVChar3 while adding either two or four additional redirecting spectrin repeats (Fig. 1d, Supplementary Videos 5 and 6 and Extended Data Fig. 2b,c). The resulting MyLOVChar4 and MyLOVChar5 designs both exhibit >80% optical modulation, on average. Individual experiments can show close to 100% optical modulation, with close to zero average gliding velocity in the dark, increasing to ~3 μ m s⁻¹ when illuminated with blue light (Fig. 1e). As expected from our design, the actin-activated ATPase rate of MyLOVChar4 is not substantially affected by light (Fig. 1g and

Supplementary Fig. 3), indicative of a gearshifting mechanism in which enzyme kinetics remain constant while mechanical output is strongly modulated via redirection of the lever arm. Although the lit velocities of the redesigned motors are reduced compared to MyLOVChar, they remain fast compared to most natural myosin classes²⁰. Thus, we succeeded in our design goal to build a molecular motor that combines high velocity with deep optical modulation.

Directional switching of gliding filaments. As shown previously¹⁰, sufficient redirection of the lever arm should yield velocity modulation that exceeds 100%, crossing the origin to produce a change in direction. For MyLOVChar variants, this would give (-)-end directed motility in dark switching to (+)-end directed motility in the light (Extended Data Fig. 2b,c). Close inspection of the gliding filament results for MyLOVChar4 and MyLOVChar5 indeed reveals some bidirectional behavior. When signed velocities are averaged over many filaments, the mean velocities are close to zero in the dark. However, a fraction of the filaments display clear (-)-end directed motion in the dark (Fig. 1e), and individual filaments can be seen to switch their direction of motion as the light is turned on (Supplementary Video 7). The stochastic behavior of individual filaments driven by small collections of these gearshifting motors thus yields distributions of velocities that extend into the (-)-end directed regime in the dark, switching to robust, rapid (+)-end directed motility in the light.

Dose-dependent response of gearshifting ensembles. A potentially useful feature of optically controlled motors is that the properties of motor ensembles may be tunable by altering the irradiance of the light signal. In isolated LOV domains, the fractional population in the lit state can be varied by changing the light intensity²¹, with steady-state populations resulting from a light-dependent photoproduct formation and a thermally driven recovery. In ensembles of optically controlled motors, the steady-state fraction of motors in the lit state thus depends on the irradiance, and the velocity of a filament or cargo driven by the ensemble may be continuously tuned to ensemble-averaged values that lie between the lit and dark extremes. We used gliding filament assays to analyze the dynamic and steady-state response of MyLOVChar4 ensembles to varying doses of blue light. As expected, at low irradiances, we found that the lit state gliding velocity depended on the blue light intensity (Fig. 1h, Extended Data Fig. 3 and Supplementary Fig. 4). We also characterized the gliding velocity transients on turning the blue light on and off. The dose-dependent rates of transition and steady-state velocities can be globally described by a simple two-state model for LOV photoswitching, in which the transition rate from the dark to the lit state depends linearly on the irradiance, while the rate of thermal decay to the dark state is fixed²¹ (Methods and Supplementary Note 1). Saturated velocities can be achieved with an irradiance of a few mW cm⁻², two to three orders of magnitude smaller than the typical intensities used in standard epifluorescence experiments on live cells²². To further test the dose-response model, we directly probed the dose-dependent conformational change of the optically controllable lever arm of the motor, using Förster resonance energy transfer (FRET) between a pair of fluorescent dyes conjugated to a minimal LOV-containing lever arm construct (Extended Data Fig. 4a,b). Bulk fluorescence measurements (Extended Data Fig. 4c-f) show a light-dependent response that saturates at similar blue light intensities to the gliding filament velocities (Extended Data Fig. 4g), as expected if the dose-dependent characteristics of the motor are dominated by the conformational changes of the LOV domain.

Directional control of processive myosins in vitro. Transport functions carried out by a number of cellular cytoskeletal motors depend on processivity—the ability of a motor to take many steps along a filament before dissociating²³. Light-activated gearshifting

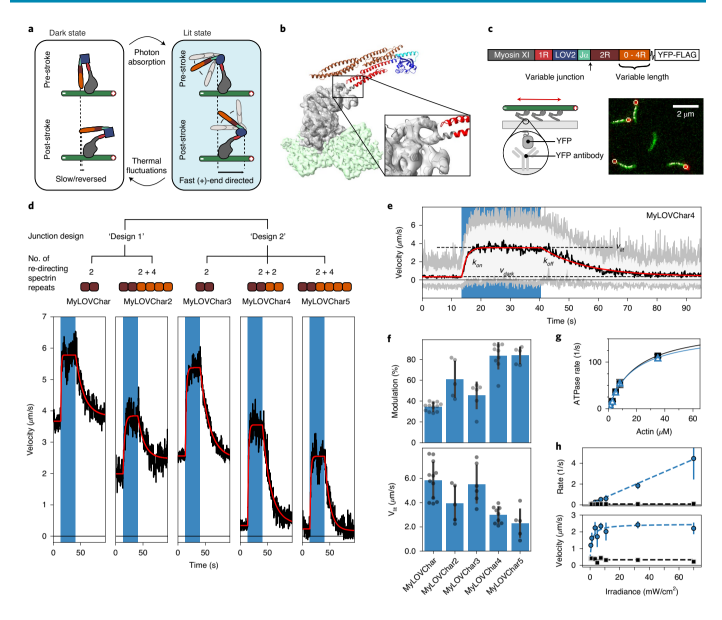


Fig. 1| Engineering monomeric myosin XI motors for high speed and strong optical response. a, Cartoon model of the mechanism of optical switching. b, Cryo-EM reconstruction of the Chara corallina myosin XI catalytic domain (apo, gray) bound to F-actin (green) and a post-stroke model of MyLOVChar (ribbon diagram) based on the cryo-EM model, with a fused lever arm structure built from crystal structure fragments using RosettaRemodel (Extended Data Figs. 1d and 2a). c, Block diagram of the molecular construct (top) and schematic (bottom left) and snapshot (bottom right) of the gliding filament assay with automated tracking overlay. d, Representative velocity traces for a series of MyLOVChar variant constructs with two different junction designs and variable lengths of the redirected lever arm. The plotted velocities are the mean directionally scored frame-to-frame velocities of all moving filaments in a video frame, averaged over three to five sequential acquisitions of the same illumination sequence on the same field of view in one assay chamber (the blue shaded areas indicate periods of blue light illumination, with irradiance of 11 mW cm⁻²). Gliding assays were performed at 2 mM ATP. Red lines are fits to the exponential rise and decay of the velocity. e, Velocity trace of MyLOVChar4 from d, in more detail. Gray and white colored bands display the envelopes of the center 96 and 72 percentiles of the distribution of frame-to-frame velocities, respectively. The model fit (red line) yields the annotated rate constants k_{on} and k_{off} and the steady-state velocities in the dark (v_{dark}) and in the presence of blue light (v_{lit}). f, Velocity modulation and lit-state velocity for the series of constructs in **d**, with each gray dot representing the result of an independent experiment analyzed as in **e**, and each bar displaying the mean ± standard deviation across the results shown (between N=5 and N=12 experiments depending on the construct). The velocity modulation is defined as $1-v_{task}/v_{lit}$ and is displayed as a percentage. g, ATPase activity of MyLOVChar4 measured with an NADH enzyme-coupled assay, with (blue data points) and without (black data points) blue light at an irradiance of 10 mW cm⁻². Lines are fits to a Michaelis-Menten expression, with $k_{cat} = 180 \, \text{s}^{-1}$ and $K_{ATPase,actin} = 20 \, \mu\text{M}$ without blue light (black line), and $k_{cat} = 160 \, \text{s}^{-1}$ and $K_{ATPase,actin} = 18 \, \mu\text{M}$ with blue light (blue line). Data are plotted as the mean of one to five enzyme velocity measurements at each actin concentration (see Supplementary Fig. 3 for details, including individual measurements and N values). Error bars display standard deviations for datasets with three or more measurements. h, Characterization of MyLOVChar4 as a function of blue light intensity (Extended Data Fig. 3). Blue data points: k_{on} (top) and v_{lit} (bottom). Black data points: k_{off} (top) and v_{dark} (bottom). The dotted lines are fits to a dose-response model, with $V_{\text{dark state}} = 0.33 \, \mu \text{m s}^{-1}$, $k_{\text{off}} = 0.079 \, \text{s}^{-1}$, $V_{\text{lit state}} = 2.5 \, \mu \text{m s}^{-1}$ and the slope of the rate dependence $m = 0.062 \, \text{s}^{-1} \, \text{mW}^{-1} \, \text{cm}^2$ (Methods). Each data point represents the mean ± standard deviation from three to five independently prepared assays (see Supplementary Fig. 4 for details); for each assay, parameters were obtained as in e from velocity traces averaged over three sequential acquisitions of the stimulation sequence.

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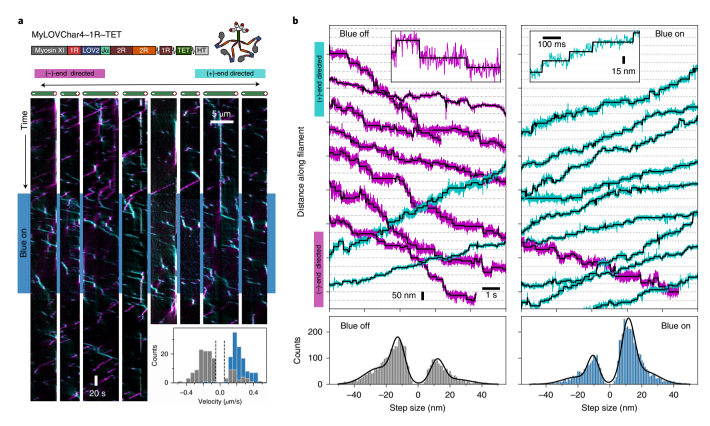


Fig. 2 | Directional control of processive myosins in vitro. a, Characterization of an optically controllable tetrameric myosin motor MyLOVChar4-1R-TET using single-molecule fluorescence. Top: block diagram and cartoon of the molecular construct. 1R-TET, a tetramerization domain with flexible linkages⁷. Main: representative kymographs displaying processive runs on four actin filaments (left and right sets of four kymographs each show the results of a first and second stimulus sequence on the same filaments). The blue band in the kymograph indicates the period when the blue light-emitting diode (LED; irradiance, 11 mW cm⁻²) was turned on. The (+)-end directed runs are colored cyan and the (-)-end directed runs are colored magenta, using an automated kymograph analysis⁴². Inset: histograms of the velocities of detected directional runs (blue, lit; gray, dark). Dashed lines indicate the minimum threshold speed used for scoring runs. The mean and standard deviation of (–)-end directed velocities in the dark is $v_{\text{dark}} = -0.18 \pm 0.08 \, \mu \text{m s}^{-1}$ (N = 138 runs), and in the light is $v_{lit} = 0.23 \pm 0.09 \, \mu m \, s^{-1}$ ($N = 123 \, runs$), with results collected over four assay chamber preparations and two different protein preparations. Data were collected at 10 μ M ATP. **b**, Gold nanoparticle tracking motor stepping traces on MyLOVChar4_{L2(+4)}-1R-TET, a variant that differs from the construct MyLOVChar4-1R-TET only by mutations to an actin-binding loop in the catalytic domain of Chara corallina myosin XI⁴³. Top: representative stepping traces with blue light off (left) and blue light on (right). Raw trajectories (colored traces, acquired at 503 Hz) are shown together with idealizations (black) based on automated step finding⁴⁴. Predominantly (+)-end directed traces are colored in cyan, while predominantly (-)-end directed traces are colored in magenta. Insets show selected traces at high resolution. Scale bars (15 nm and 100 ms) apply to both insets. Data were collected at $5\,\mu\text{M}$ ATP. Bottom: step size distributions in the annotated light condition. Distributions of scored step sizes (blue off, N = 4,364; blue on, N = 4,809) are shown together with fits (black lines) of sums of Gaussian distributions. Peak locations \pm standard errors are as follows. Blue off, -24.7 ± 0.4 nm, -12.4 ± 0.3 nm, $+12.1 \pm 0.4$ nm and $+24.0 \pm 0.9$ nm; blue on, -20.9 ± 0.6 nm, -10.4 ± 0.3 nm, $+11.4 \pm 0.2$ nm and $+22.1 \pm 0.4$ nm.

has not previously been combined with processive motion, although controllable bidirectional processive myosins have been engineered that respond to metal ions^{6,7} or oligonucleotide strands¹⁵. We designed a controllable processive motor based on MyLOVChar4 by adopting a strategy of forced tetramerization in combination with flexible linkers (Fig. 2a). In previous work, this strategy was found to be successful for building highly processive motors from engineered monomers, and was applied to a fixed-gear motor based on the *Chara* myosin XI catalytic domain⁷.

We characterized the performance of MyLOVChar4~1R~TET (Fig. 2a and Supplementary Video 8) using in vitro single-molecule tracking, where single fluorescently labeled motor complexes are imaged translocating along actin filaments immobilized on a glass coverslip. In the absence of blue light (top and bottom part of the kymograph, Fig. 2a), the majority of motors move toward the actin (–)-end. A small fraction of runs are (+)-end directed, revealing the heterogeneity of motor function at the single-molecule level. When the blue light is turned on, almost all motors become (+)-end

directed (Fig. 2a kymograph, center region with blue band). This behavior is also seen in a histogram (Fig. 2a, inset) of several hundred motor runs, with almost exclusively (+)-end directed motion in the presence of blue light and predominantly (-)-end directed motion in the absence of blue light. To probe the stepping behavior of bidirectional motors, we performed high-resolution gold nanoparticle tracking⁷ using a variant of MyLOVChar4~1R~TET (Fig. 2b and Supplementary Video 9). Nanoparticles also show processive light-dependent bidirectional motion, with discrete steps identified using automated change-point detection (insets, Fig. 2b and Supplementary Fig. 5). As seen previously for unidirectional tetrameric myosins incorporating flexible spacer elements⁷, the step size distributions are broad, with frequent backsteps. Steps are predominantly (+)-end directed in the presence of blue light and predominantly (-)-end directed in the absence of blue light (bottom, Fig. 2b and Supplementary Fig. 6), with distributions peaked at similar magnitudes and opposite signs in the two conditions (dominants peaks in Fig. 2b; blue off, $-11.4 \,\mathrm{nm}$; blue on, $12.4 \,\mathrm{nm}$).

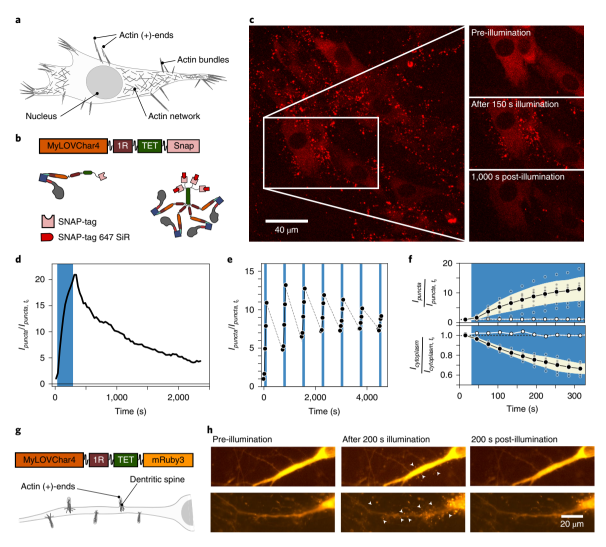


Fig. 3 | Optically controlled localization of myosin motors in live cells. a, Simplified schematic of a cell with different actin architectures at various locations in the cell. b, Block diagram and cartoon of the molecular construct imaged in live cells. c, Maximum intensity projection of a three-dimensional (3D) confocal stack of fluorescence images (excitation at 639 nm) of a sheet of mouse fibroblasts in culture immediately after 150 s of stimulation with blue light, displaying fluorescent puncta concentrated around the periphery of the cell. Insets show a region of the sheet before and after blue light stimulation. Images are representative of a set of seven independent experiments with comparable blue light stimulation protocols (Supplementary Fig. 7). d-f, Fluorescence intensity contained in puncta, I_{punctar} , normalized to the total intensity in the image and plotted after further normalization to the value at the start of the series, Ipuncta.tn. The blue bands indicate the period of blue light stimulation using a blue laser line in the scanning confocal microscope. d, Intensity in puncta calculated for the image series on the cells in c. e, Intensity in puncta over seven stimulation cycles of 2 min each, separated by 40-min wait times. In each stimulation cycle, cells were imaged for five frames, with one frame without blue light preceding four frames with blue light on. Gray dashed lines are visualization guides. f, Time traces of averaged responses (black data points) upon stimulation with blue light, from seven independent sample preparations (gray data points). Black data points represent the means of the seven measurements. Error bands indicate the standard deviation of the measurements. White data points represent the response in the absence of blue light. Top: fluorescence intensity contained in puncta; the maximum level of I_{puncta,I_0} after 300 s of stimulation was 11 ± 4. Bottom: fluorescence of cytoplasm. **g,h**, Characterization of MyLOVChar4~1R-TET~mRuby3 in hippocampal neurons in culture: block diagram of molecular construct and illustration of actin architecture in neurons (g) and wide-field fluorescent images of MyLOVChar4-1R-TET-mRuby3 before, during and after stimulation with blue light (h) (Supplementary Video 13). Arrows in h highlight puncta as expected for motor accumulation in dendritic spines. Data are representative of an experiment replicated twice on different days, with 13 cells.

As expected, characteristic dwell times between detected steps do not substantially change upon illumination (Supplementary Fig. 6; blue off, $0.235\pm0.004\,s$; blue on, $0.246\pm0.004\,s$ at $5\,\mu M$ ATP). Using both fluorescence and gold nanoparticle tracking, we have thus shown that engineered processive myosins can switch their direction of motion under the influence of an optical signal in a reconstituted system with purified proteins.

Optical control of motor localization in living cells. To satisfy our final performance criterion, we asked whether

MyLOVChar4~1R~TET retained motility and optically controlled directionality in living cells. We characterized motor function in fibroblasts and in hippocampal neurons. Actin filament architectures within the cell include branched networks of cortical actin close to the cell membrane and polarized actin bundles in protrusions around the periphery of the cells, with actin (+)-ends at the protrusion tips²⁴ (Fig. 3a). For experiments in fibroblasts, we expressed MyLOVChar4~1R~TET as a SNAP-tag fusion, and imaged the motors labeled with a red fluorescent dye (Fig. 3b and Methods). Before optical stimulation, motors were distributed throughout the

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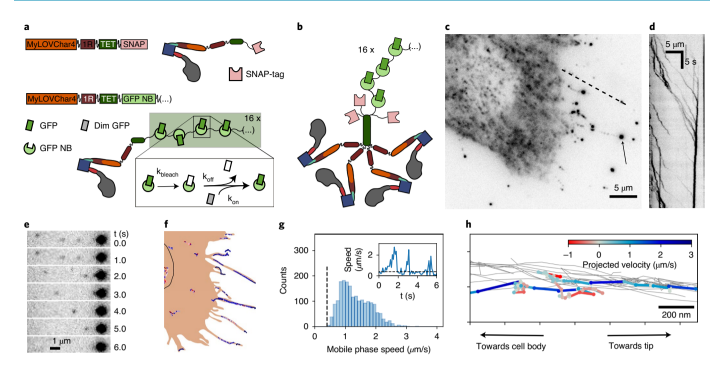


Fig. 4 | Single-molecule imaging of engineered motor constructs in live cells. a, Design of molecular constructs for single-molecule tracking with high spatiotemporal resolution in cells. The block diagram and cartoons show two MyLOVChar4 constructs, C-terminally tagged with either a SNAP-tag (MyLOVChar4-1R-TET-SNAP) or a 16X ArrayG scaffold of GFP nanobodies (MyLOVChar4-1R-TET-ArrayG_{16X}). Dim monomeric wild-type GFP (mwtGFP, gray rectangles) becomes brighter (green rectangles) upon binding to the scaffold, and reversible association allows exchange of bleached mwtGFP (white rectangles) with unbleached protein from solution. **b**, Cartoon of the tetrameric motor complex imaged in cells overexpressing the non-visible species MyLOVChar4-1R-TET-SNAP, a minor population of MyLOVChar4-1R-TET-ArrayG_{16X} and mwtGFP. **c**, Fluorescence image of a cell excited with 488 nm laser excitation. Arrow: bright fluorescence spot that is a terminal point of the observed tracks. **d**, Kymograph along the dashed line in **c**. **e**,**f**, Single-molecule tracking of motors in cells. The image series in **e** is taken from a region of the cell in **c**, with single motors moving toward, and merging with, a bright spot like the one indicated in **c**. Motor trajectories identified for the cell in **c** are shown in **f**, overlaid on a schematic outline of the cell. The black outline shows the estimated location of the nucleus. Similar results to those presented in **c**-**f** were obtained on two independent biological replicates, across a total of 41 cells contained in 39 image acquisition series. **g**, Velocity histogram from all 41 cells, for all extracted directionally persistent tracks with a minimum length of 20 frames, tabulating mobile runs in which the speed exceeds 0.4 μm s⁻¹ for four frames or longer (*N*=1,946 runs). The mean and mode of the distribution are 1.3 μm s⁻¹ and 1.0 μm s⁻¹, respectively. Inset: velocity trace with definition of the mobile phases. **h**, Detail of motor trajectories in a cellular protrusion. Gray thin line

cytoplasm (Fig. 3c and Supplementary Video 10). Within 30s of illumination with blue light, distinct puncta became visible around the periphery of the cell, increasing in number and intensity over a 150-s stimulation period, and decaying after the removal of blue light on a longer timescale of 100-1,000s (Fig. 3c,d and Supplementary Fig. 7). We attribute these puncta to motors accumulating at the tips of cellular protrusions, after switching to (+)-end directed motion at the start of optical stimulation and translocating out to the (+)-ends of polarized actin structures. To further support this attribution, we simultaneously imaged the fluorescence of motors and the contours of the cell with differential interference contrast microscopy, and found that puncta co-localize with tips of protrusions (Supplementary Video 11). In the absence of blue light, the strongly reduced number and intensity of puncta are consistent with directional motion of a dominant fraction of motors toward the actin (-)-end in the dark. Translocation can be repeated across multiple cycles of illumination and recovery (Fig. 3e), with residual persistence of some puncta in the dark, leading to reduced effects over successive cycles. The population of motors in puncta depends monotonically on blue light intensity at low irradiances (Supplementary Fig. 7), and strong effects are seen at irradiances on the order of 1 mW cm⁻², comparable to levels commonly used for optogenetic experiments²⁵. The cytoplasmic intensity decreases concomitantly with the appearance of puncta (Fig. 3f and Supplementary Video 12), indicating that a substantial fraction of the total population of motors within the cell can be sequestered in cellular protrusions. Experiments in cultured neurons also revealed a rapid formation of fluorescent motor puncta around the periphery of neurites upon stimulation with blue light (Fig. 3g,h and Supplementary Video 13), consistent with motors accumulating at the tips of dendritic spines, where actin filaments are polarized with (+)-ends outward²⁶. Taken together, our results show that we can optically control the localization of MyLOVChar4~1R~TET motors in cells.

Single-molecule tracking in living cells. To directly measure the dynamics and motor properties of individual engineered myosins in cells, we performed high-resolution single-molecule tracking (Fig. 4 and Supplementary Video 14). Earlier reports of single-molecule tracking of motors in cells have used quantum dots²⁷, individual fluorescent protein (green fluorescent protein, GFP) tags²⁸, GFP-tagged peroxisomes²⁹, and scaffold arrays that tightly bind multiple fluorescent proteins³⁰. Here we use a 16-copy version of the fluorogenic array tag ArrayG³¹, where up to 16 monomeric wild-type GFPs (mwtGFPs) can bind to a scaffold of flexibly enchained camelid nanobodies. The mwtGFPs bind reversibly, so the effect of bleaching can be diminished by exchange with a pool

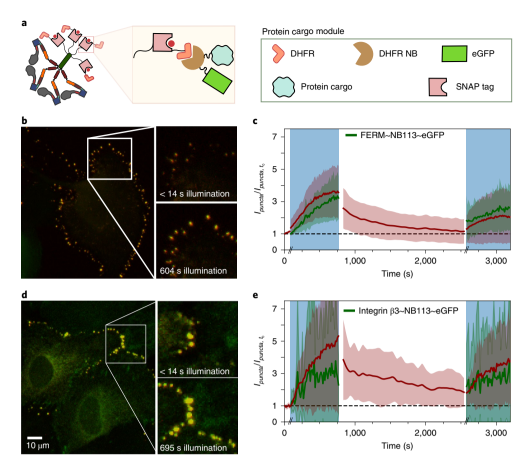


Fig. 5 | Optically controlled localization of motors and molecular cargos in live cells. a, Schematic of MyLOVChar4-1R-TET-DFHR interacting with a protein cargo fused to a DFHR nanobody (NB113). **b-e**, Results from cells expressing MyLOVChar4-1R-TET-DFHR and FERM-NB113-eGFP (**b,c**) and cells expressing MyLOVChar4-1R-TET-DFHR and integrin β3-NB113-eGFP (**d,e**). Merged two-channel fluorescence images (maximum intensity projections of confocal *z*-stacks; red, motor; green, cargo) from sheets of MCF10A cells in culture are shown in **b** and **d**, imaged with simultaneous 639 nm and 488 nm excitation. Images are representative of a set of 10 (**b**) and 5 (**d**) independent measurements, on 12 and 8 cells, respectively. The fluorescence intensity contained in puncta is shown in **c** and **e**, normalized to values at the start of the imaging sequence, for motor (red) and cargo (green) channels. Results are averaged over individual time series (Supplementary Figs. 9 and 10) representing 12 (**c**) and 8 (**e**) cells, from two biological replicates in each case. Error bands are standard deviations of the measurements. The time axes include approximate delays at transitions between the illumination phases (indicated by the double dashes on the axes), which for these experiments were collected as separate videos.

of unbleached mwtGFPs in the cytoplasm. mwtGFP fluorescence increases by ~26-fold on binding to the array, which reduces the imaging background (Fig. 4a). To favor robust tetramer (Fig. 4b) formation while keeping the concentration of the Array G_{16X} tag low enough for single-molecule imaging, we co-expressed low levels of Array G_{16X} -tagged motors with an excess of unlabeled SNAP-tagged motors (Methods).

Single-molecule imaging of Array G_{16X} -tagged MyLOV4~1R~TET was performed under blue laser excitation, which both activates the LOV2 domain and excites the GFP fluorophore. Under these conditions, we expect the motors to be predominantly (+)-end directed. As expected, fluorescent spots around the periphery of the cell (Fig. 4c) are seen streaming away from the cell body along selected paths (Supplementary Video 14), as shown in a kymograph (Fig. 4d) and image series (Fig. 4e) along selected paths. Single-molecule trajectories often terminate in a brightly labeled endpoint that increases in intensity over time (Fig. 4e and Supplementary Video 15). Supported by our observations of the cellular morphology and ensemble motor localization (Fig. 3 and Supplementary Videos 10 and 11), we conclude these trajectories correspond to motors traveling toward the actin (+)-end on cellular protrusions containing polarized actin bundles.

To measure motor velocities in cells, we extracted high-resolution traces from the fluorescence imaging series using a tracking algorithm optimized for high spot densities³². Extracted traces for an example cell (Fig. 4c) show many traces around the cell periphery, a set of traces in the region below the nucleus and a few isolated traces in between (Fig. 4f). The imaging conditions in our experiment are most favorable for the detection of processive runs in thin cellular protrusions, because of the dense, polarized actin structure and lower fluorescence background. The motors predominantly travel away from the cell body, moving in intermittent bursts of high velocity separated by much slower phases (inset, Fig. 4g and Supplementary Fig. 9). We analyzed the frame-to-frame speeds of motors tracked in 53 cells and found a broad distribution of mobile phase speeds with a peak at ~1 µm s⁻¹ (Fig. 4g). Distinct trajectories can be resolved within sets of clustered traces around the cell periphery (Fig. 4h), revealing the spatial extent of the protrusions that motors are able to explore. Short directional reversals can be seen for some traces (Fig. 4h and Supplementary Fig. 9), consistent with motors that temporarily become (–)-end directed. Taken together, these results show that optically activated motors can efficiently traverse across various cellular actin structures, with velocities in the range of micrometers per second.

Controllable transport of molecular cargos. Having satisfied our performance criteria for controllable processive molecular motors that function in living cells, we set out to directly test whether the engineered motors could be harnessed for optically controlled cargo transport (Fig. 5). Native myosins are responsible for transporting diverse molecular and vesicular cargos³³. We chose to test two distinct classes of molecular cargos: the talin FERM domain³⁴ (a cytosolic protein) and integrin β3 (an integral membrane protein). Previous work has shown that fusion of a myosin to a protein of interest or nanobody can be sufficient for localizing a binding partner into cellular protrusions³⁵. We sought to couple controllable motors to protein cargos via a nanobody interaction³¹ (Methods and Fig. 5a), and co-expressed motor and cargo constructs in epithelial cells. In these experiments, the tetrameric motor can be visualized with a SNAP-tag fluorophore excited by a red laser, while blue excitation is used to both visualize the fluorescent cargo and switch the motor directionality. Upon stimulation with blue light, both the cytoplasmic FERM domain (Fig. 5b,c, Supplementary Fig. 10 and Supplementary Videos 16 and 17) and the transmembrane integrin β3 (Fig. 5d,e, Supplementary Fig. 11 and Supplementary Videos 18 and 19) are enriched in puncta around the cell periphery, closely following motor enrichment. The localization of motors largely decays within several hundred seconds when blue light is removed, while a second stimulus reveals a more persistent localization of the cargos that increases again during the illumination period (Fig. 5c,e). Taken together, our observations show that an engineered myosin motor with optically controllable directionality can localize distinct classes of molecular cargos to cellular protrusions.

Discussion

We have presented a set of engineered myosins with optically controllable velocities and directionalities, with speeds matching the range of natural biological motors. Deviations in modulation depths from the predictions of idealized structural models (Extended Data Fig. 2a-c) can be overcome by extending redirected lever arms and optimizing engineered junctions. Monomeric motors can be assembled into controllable processive complexes and used for light-dependent intracellular transport of cargos. This strategy complements approaches based on light-dependent cargo recruitment¹¹⁻¹⁴ while avoiding the challenges associated with non-productive accumulation of the motor pool at target locations³⁶ and allowing for the use of tight constitutive cargo-binding interactions that can resist the forces generated in cellular contexts. Light-activated gearshifting with photoswitchable cargo binding at orthogonal wavelengths³⁷ could also provide additional levels of control. Fast monomeric non-processive gearshifting motors may be used to introduce programmability into devices that rely on gliding filament propulsion³⁸, or spatial control in active fluids composed of propelled filaments³⁹. The multimeric motors here could also be used for spatiotemporal control of active matter systems⁴⁰, and similar design principles may be used to control the mechanical output of filamentous myosins to enable control of cellular contractility. Further extension of the gearshifting mechanism to microtubule-based motors with swinging lever arm mechanisms such as kinesin-1410 and dynein41 would then complete a toolbox of optically controlled motors that can be used to probe the relationships between nanoscale motility and complex cellular functions.

Online content

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Methods

Molecular constructs for in vitro experiments. Myosin constructs were assembled from DNA fragments encoding Chara corallina myosin XI (codons 1-746), Nicotiana tabacum myosin XI (codons 1-738), Dictyostelium discoideum α-actinin (codons 266–502), Avena sativa phot1 (codons 404–543) and Gallus gallus α-spectrin (codons 1663-2090) and cloned into the expression plasmid pBiEx-1 (Novagen). Constructs encoding the monomeric motors MyLOVChar and MyLOVChar1-5 include codons for a C-terminal flexible linker, enhanced yellow fluorescent protein (YFP) and FLAG tag (DYKDDDDK) as in previous work¹⁰. Constructs for the tetrameric motors MyLOVChar4~1R~TET and NM11CD738_2R~1R~TET include codons for a C-terminal engineered leucine zipper variant pLI⁴⁵, a spacer consisting of *Dictyostelium discoideum* α-actinin codons 266-388 flanked by flexible linkers, a Halotag protein sequence (Promega) and a FLAG tag (DYKDDDDK) as in previous work⁷. Motor proteins were expressed by direct transfection in Sf9 insect cells and purified using anti-FLAG resin (Sigma-Aldrich A2220), as previously described^{7,46}. Before transfection, the cells were cultured in serum-free medium without antibiotics. Myosin proteins used for in vitro assays were eluted in a glycerol-containing storage buffer (25 mM KCl, 2 mM MgCl₂, 5 mM EGTA, 25 mM imidazole pH 7.5, 0.1 mM ATP, 1 µg ml⁻¹ aprotinin, 1 µg ml⁻¹ leupeptin, 2 mM DTT and 55% glycerol by volume), flash-frozen in small aliquots within hours of the completed elution, and stored at -80 °C until used in assays. The myosin construct characterized by electron microscopy (CM11_{L2+4}CD746) was eluted in a glycerol-free storage buffer (25 mM KCl, 2 mM MgCl₂, 5 mM EGTA, 25 mM imidazole pH 7.5, 0.1 mM ATP and 2 mM DTT) and concentrated using 10 kDa Amicon Ultra Millipore columns before flash freezing. For details of the constructs see Supplementary Fig. 2.

The minimal lever arm construct SNAP-1R-LOV-1R-HT for FRET measurements was assembled with DNA fragments coding for a section of the lever arm from MyLOVChar4 as described above and cloned into the bacterial expression vector pET24a. The construct also includes codons for an N-terminal SNAP tag and a C-terminal Halotag and His-Tag. The construct was expressed in Escherichia coli B21 (DE3) by induction with 0.5 mM isopropyl-β-D-thiogalactoside at 16 °C in 2xYT broth for 20 h. Cells were collected by centrifugation and frozen. Bacterial pellets were thawed, lysed with hen egg lysozyme (1.7 mg ml⁻¹) and by mechanical lysis with a sonicator, and purified by metal ion affinity (Ni-NTA column, HisPur resin, Thermo Scientific) and size-exclusion chromatography (Superdex 200/10/300 GL). Lysis buffer consisted of 50 mM Tris (pH 7.5), 500 mM NaCl and 1.7 mM PMSF. Wash buffer consisted of PBS with 50 mM imidazole (pH 7.5). Elution buffer consisted of PBS with 500 mM imidazole (pH 7.5). Following the metal affinity columns, proteins were dialyzed overnight into PBS with 0.5 mM DTT and concentrated using 3 kDa Amicon Ultra Millipore columns. Concentrations were measured by absorbance at 280 nm using a Nanodrop spectrophotometer (Thermo Scientific). For details of the construct see Supplementary Fig. 2.

Cryo-electron microscopy on Chara myosin XI. Actin purification. Chicken muscle actin was purified as previously described and stored in G-Mg buffer (2 mM Tris-Hcl pH 8.0, 0.5 mM DTT, 0.2 mM MgCl₂ and 0.01% NaN₃). F-actin was prepared by polymerizing 10 μ M monomeric actin in KMEI buffer (50 mM KCl, 1 mM MgCl₂, 1 mM EGTA and 10 mM imidazole pH 7.0) overnight at 4 °C. Phalloidin (20 μ M) was added to the F-actin 30 min before grid preparation.

Grid preparation and cryo-electron microscopy image acquisition. Before vitrification, 10 U ml⁻¹ apyrase was added to 18.5 μM myosin XI to deplete nucleotide. Phalloidin-bound F-actin (3 µl), diluted to 0.6 µM, was applied to a glow-discharged C-flat 1.2/1.3 holey-carbon Au 300 mesh grid (Electron Microscopy Sciences) in a Leica GP plunge freezer at 25 °C. After 60 s, 3 μl of myosin XI was applied and incubated for 60 s, then 3 µl of the mixture solution was removed and another 3 µl of myosin XI applied. After an additional 60 s of incubation, 3 µl of the mixture solution was removed and the grid was blotted from the back with Whatman no. 5 filter paper for 5s and flash-frozen in liquid ethane. Data were collected on an FEI Titan Krios system (New York Structural Biology Center) operating at 300 kV and equipped with a Gatan K2-summit detector and a Cs-corrector using counting mode. Image stacks were recorded with Leginon software at a nominal magnification of ×105,000, corresponding to a pixel size of 1.096 Å. Each exposure was fractionated across 50 frames with a total electron dose of 67.12 e⁻ per Å² (1.34 e⁻ per Å² per frame) and a total exposure time of 10 s. Images were recorded with defocus values ranging from -1 to $-4 \mu m$ underfocus. In total, 714 movie stacks were acquired.

Cryo-electron microscopy image processing. Data processing was performed as described in ref. 48. Briefly, image processing was carried out using the Iterative Helical Real Space Refinement (IHRSR) protocol as implemented in the RELION 3.0 pipeline 49, utilizing RELION functions unless otherwise noted. Movie stacks were motion-corrected and dose-weighted with MotionCor2 (https://emcore. ucsf.edu/ucsf-software) using 5×5 patches. Contrast transfer function (CTF) estimation was performed with CTFFIND4 (https://grigoriefflab.umassmed. edu/ctffind4). Filaments were auto-picked and split into overlapping segments with a step size of 81 Å, corresponding to three actin protomers. A total of 72,336

segments were extracted with a box size of 512 pixels. After 2D classification and sorting, 45,779 segments were selected and subjected to 3D classification initialized with the helical parameters of 27 Å rise and –166.7° twist. The cryo-EM map of myosin VI bound to F-actin (EMD-7116) was low-pass-filtered to 30 Å to serve as an initial model. The 3D auto-refinement and post-processing using a 3D mask with a length of 30% of the box size resulted in a 5.7 Å map. CTF refinement, Bayesian polishing and another round of 3D refinement improved the resolution to 4.3 Å based on the Fourier shell correlation (FSC) 0.143 criterion. Final helical parameters converged to a rise of 27.45 Å and –167.11°. Local resolution was estimated using RELION 3.0. Figures were generated in UCSF Chimera and ChimeraX⁵⁰

Model building and structure refinement. The initial model of myosin XI was generated with I-TASSER (https://zhanglab.ccmb.med.umich.edu/I-TASSER). The myosin model and a published actin model (PDB 6BNO) were docked into the EM density by rigid body fitting in UCSF Chimera. This initial actomyosin model was then rebuilt and refined as described in ref. 48. Briefly, 100 models were generated and the lowest energy model was manually adjusted in Coot (https://www2.mrc-lmb.cam.ac.uk/personal/pemsley/coot) and refined using Phenix.real_space. refine in the Phenix suite 51. As the region of the cryo-EM map corresponding to myosin XI was of limited resolution (5–8 Å, Extended Data Fig. 1c), and largely lacked well-defined side-chain density, the myosin chain of the atomic model was truncated to poly-alanine before deposition.

Structural modeling of optically controllable motor constructs. We generated idealized pre- and post-stroke structures for MyLOVChar-MyLOVChar5 using flexible backbone modeling with RosettaRemodel⁵² to model engineered junctions between protein domains. For modeling the catalytic domain, in post-stroke models we used the rigor structure of Chara corallina myosin XI obtained in this work (see previous section), and in pre-stroke models we used a previously reported structure of human myosin Vc (PDB 5HMP)53, a myosin isoform related to plant myosin XI. Engineered lever arms were constructed from available crystal structures for Dictyostelium discoideum α-actinin (PDB 1g8xA residues 766-1002), Avena sativa phot1 (PDB 2V0U) and (for MyLOVChar2, MyLOVChar4 and MyLOVChar5) Gallus gallus α-spectrin (PDB 1U4Q). Models of the engineered constructs were obtained by remodeling four to eight residues on either side of each engineered junction as α -helical structures, followed by a relaxation protocol on the complete lever arm sequence. See Supplementary Fig. 1 for examples of the RosettaRemodel flags and blueprint files with the instructions for modeling of the junctions. We gathered 500 candidate structures for each model, and removed structures that clashed with the actin filament (which was not included in the initial computation). To obtain the idealized predicted power strokes of the motors in the light and dark states, we aligned the 50 top-scoring structures to a subregion (corresponding approximately to the lower 50-kDa domain L50 of myosin) of the cryo-electron microscopy reconstruction of rigor Chara corallina myosin XI bound to actin, and extracted the mean projected stroke along the filament of the end of the lever arm (for the dark state) or the last rigid residue on the LOV2 domain, taken as G516 (for the lit state).

Gliding filament motility assays. Monomeric engineered myosin constructs were characterized using a dual-labeled gliding filament assay as described in ref. 10 , with minor modifications to buffers and imaging conditions, as described below. Flow channels were assembled with microscope coverslips attached to a cover slide with double-sided Scotch tape. Channels were $\sim\!2$ mm wide and had a volume of $\sim\!4$ –5 µl. Before assembly of the channel, the coverslips were cleaned by 5 min of plasma cleaning, and spin-coated with 0.1% nitrocellulose (Ladd Research 10800) for 30 s at 3,000 r.p.m., following by 30 s at 6,000 r.p.m., with ramps of 5s.

Assay buffer consisted of 25 mM Tris (pH7.5), 25 mM KCl, 1 mM EGTA, 2 mM MgCl₂ and 10 mM DTT. Blocking buffer consisted of assay buffer supplemented with 2 mg ml⁻¹ BSA (Sigma-Aldrich). Imaging buffer consisted of blocking buffer supplemented with an oxygen-scavenging system (0.2 mg ml⁻¹ glucose oxidase, 0.36 mg ml⁻¹ catalase and 0.4% wt/vol glucose) and an ATP regeneration system (creatine phosphokinase at 0.95 μ g ml⁻¹ and 1 mM phosphocreatine). The assay conditions differ from those in ref. 10 due to the complete omission of imidazole, which even at the low levels present in the previous study can be sufficient to speed up the dark state recovery of the LOV domain by several fold 54 .

Flow channels were prepared in six steps: (1) flow in 3.5 μ l of GFP antibody (Millipore Sigma MAB3580) diluted threefold from stock in PBS, followed by an incubation period of 2 min; (2) flow in 30 μ l of blocking buffer, followed by an incubation period of 2 min; (3) flow in 10 μ l of GFP-tagged monomeric myosin motor (diluted from stock in assay buffer to 100–400 nM), followed by an incubation period of 2 min; (4) wash with 30 μ l blocking buffer; (5) flow in 30 μ l of polarity-labeled actin by gravity flow and incubation for 3 min—actin was freshly diluted 1,000-fold to 4,000-fold from stock into assay buffer for each channel, in a two-step dilution, gently tapping the Eppendorf tube between steps to mix; (6) flow in imaging buffer. Before imaging, the channels were sealed with vacuum grease.

Channels were imaged with total internal reflection fluorescence (TIRF) excitation, with a 1.49 NA, $\times 100$ objective (Nikon), excited with a 532 nm laser (Coherent Sapphire, 30 μW at the back aperture) and a 633 nm laser

(Blue Sky Research, 30-80 μW at the back aperture). Fluorescence from tetramethylrhodamine (TMR)/Cy5-labeled actin filaments was recorded on an electron-multiplying charge-coupled device (EMCCD) camera (Andor iXon DU-897E-CS0-#BV) at maximum gain. A light-emitting diode (LED) light source (Thorlabs, M470L3, center wavelength of 470 nm) provided continuous-wave blue light stimulation via the TIRF illuminator arm (Nikon T-FL-TIRF2). The blue light intensity was controlled using a voltage-driven modulating input on the LED current driver and synchronized to the camera acquisition with custom-written LabVIEW software. For most assays (Fig. 1d-f), 800-frame videos were acquired at a frame rate of 7.4 Hz (0.136 s per frame), with blue light on (irradiance of 11 mW cm⁻²) in frames 100-300. For Fig. 1h we acquired videos of 550, 650 and 750 frames, with blue light turned on during frames 50-150, 50-250 and 50-350, respectively (Extended Data Fig. 3), with the 100-frame blue light stimulation period for the highest irradiances (32 and 70 mW cm⁻²), the 200-frame stimulation period for intermediate irradiances (3.4, 4.8, 7.3 and 11 mW cm⁻²) and the 300-frame stimulation period for the lowest irradiances (0.5 and 1.2 mW cm⁻²).

Gliding filament motility analysis. Gliding filament assays were analyzed to extract the frame-to-frame velocity of directionally scored filaments using a custom workflow building on the FAST software for gliding filament motility analysis of Aksel and colleagues⁵⁵. Two-color videos with the registered images of the filaments in a green fluorescence channel and the filament ends in a red fluorescence channel were generated from the raw data using custom-written scripts in MATLAB, with image registration between the channels based on cross-correlation using a discrete Fourier transform algorithm. We used custom-written Jython scripts to automate the use of ImageJ plugins for filament detection (Ridge Detection plugin version 1.4⁵⁶, implementing the ridge detection algorithm of Steger) and tracking of polarity labels (TrackMate plugin⁵⁷). Motility was then analyzed using Python software modified from FAST⁵⁵ and incorporating stuck filament rejection, selection of polarity-marked filaments, and directionality determination.

We collected sets of signed trajectories over multiple videos (3–5 repeats of the same light stimulus sequence acquired on the same field of view in a single sample chamber, with no additional waiting time between repeats) and computed frame-to-frame filament velocities, which were averaged to obtain velocity traces as in Fig. 1d. The average number of filament velocities combined to obtain each time point in each velocity trace was $<\!N\!>=\!51$ for the dataset represented in Fig. 1d and compiled in Fig. 1f. The parameter value for the velocity in the dark before light stimulus, $v_{\rm dark,i}$, was calculated as the mean of the data points before blue light was turned on. With $v_{\rm dark,i}$ fixed, we globally fitted a single exponential rise followed by an exponential decay using a nonlinear least-squares fitting procedure (Scipy.optimize curve_fit). This fit provides the rates $k_{\rm on}$ and $k_{\rm off}$, the lit velocity $v_{\rm lat}$ and the final velocity $v_{\rm dark,i}$. In Fig. 1e,h, the reported dark velocity $v_{\rm dark}$ is taken as $(v_{\rm dark,i}+v_{\rm dark,i})/2$.

Gliding filament dose–response modeling. We modeled the dose response for MyLOVChar4 (Fig. 1h) as a two-state system with a population of motors in the dark state, $\varphi_{\rm dark\, state}$, and a population of motors in the lit state, $\varphi_{\rm lit\, state}$, and corresponding characteristic velocities, $V_{\rm dark\, state}$ and $V_{\rm lit\, state}$. In this two-state model, the lit state thermally decays into the dark state with a constant rate $k_{\rm lit \to dark \to lit}$. Transition from the dark state into the lit state occurs with a rate $k_{\rm dark \to lit}$ that depends linearly on irradiance, as

$$k_{\text{dark} \rightarrow \text{lit}} = mI$$

where I is the irradiance of the light and m the slope. The steady-state fraction of motors in the lit state is then

$$\varphi(I) = \frac{\varphi_{\rm lit\; state}}{\varphi_{\rm dark\; state} + \varphi_{\rm lit\; state}} = \frac{mI}{k_{\rm lit \rightarrow dark} + mI}$$

Assuming a simplified model for filament gliding that neglects strain-dependent effects (Supplementary Note 1), this results in a dose-dependent steady-state velocity:

$$\begin{split} V(I) &= (1 - \varphi(I)) V_{\text{dark state}} + \varphi(I) V_{\text{lit state}} \\ &= V_{\text{dark state}} + (V_{\text{lit state}} - V_{\text{dark state}}) \varphi(I) \end{split}$$

The parameters of this two-state model may be related to the parameters fit to the velocity transients (see above) after noting that $k_{\rm off}$ corresponds directly to the decay rate of the lit state population, $k_{\rm off} = k_{\rm lit-dark}$, while the measured transient rate $k_{\rm on}$ reflects the relaxation process of the two-state system, $k_{\rm on} = k_{\rm lit-dark} + k_{\rm dark-lit}$

Steady-state ATPase assay. The steady-state ATPase activity was measured using the NADH coupled assay as described in ref. 58 , but instead of the imidazole buffer used in ref. 58 , we used a Tris-based assay buffer as for the gliding motility assay (see above), supplemented with 1 mg ml $^{-1}$ BSA. In addition, we implemented the capability to illuminate the sample with blue light. Briefly, assays were performed with 14nM of MyLOVChar4 motors at 23 °C, monitoring NADH absorption at a wavelength of 340 nm in a spectrophotometer (Agilent Cary UV–vis 6000). Assays

were performed with $150\,\mu l$ of sample in a quartz cuvette with path length of 1 cm and nominal volume of $100\,\mu l$ (Starna Cells 16.100-Q-10/Z15), using a microscope coverslip as a translucent lid. Samples could be illuminated from the top of the cuvette (perpendicular to the light path of the 340 nm absorption measurement) with a 12-mm-diameter collimated beam (fiber collimator Thorlabs F950-FC-A) from a fiber-coupled light source (Thorlabs M470F3, controlled by voltage modulation with a function generator, Agilent 33220A). The blue light irradiance was $10\,\mathrm{mW\,cm^{-2}}$. In each measurement at a given actin concentration we measured slopes of decreasing absorbance during one or more cycles of a period without blue light, followed by a period with blue light. Illumination periods varied from 30 s to 2 min, for the highest and lowest actin concentration, respectively.

FRET assays on the minimal lever arm construct. We performed FRET measurements in solution on the minimal lever arm construct Snap~1R-LOV-1R~HT at 100 nM, either singly labeled (with Snap Alexa 647-SiR or with Halotag TMR) or doubly labeled (Snap Alexa 647-SiR and Halotag TMR). Labeling was done on flash-frozen aliquots of the purified protein (see section 'Molecular constructs for in vitro experiments') in a single step rotating at 4 °C for 3.5 h in PBS buffer. We used a fivefold molar excess of the ligands, which were removed after the labeling by overnight dialysis into Tris assay buffer (25 mM Tris, 25 mM KCl, 1 mM EGTA, 1 mM DTT and 2 mM MgCl_2), with two buffer exchanges and >500:1 sample-to-buffer volume ratio. Labeled proteins were flash-frozen in aliquots for single-use experiments.

Fluorescence emission spectra and time-resolved measurements were collected on a fluorimeter (Horiba Fluorolog 3) with a 450 W xenon lamp light source. Samples were contained in a quartz cuvette with a nominal volume of $40\,\mu$ (Starna Cells 16.40F-Q-10/Z15). Samples could be stimulated with blue light during fluorescence emission experiments through the exit window in the cuvette, illuminated through a hole in the front panel of the fluorimeter by a fiber-coupled LED light source as used in ATPase activity assays. The intensity of the 12-mm collimated beam was further concentrated with a 50-mm lens to slightly overfill the $2\times2\,\mathrm{mm}$ exit window. Blue light irradiance was controlled by voltage modulation with a function generator as used in the ATPase activity assay, and insertion of appropriate optical density filters. The blue light was spectrally filtered with a bandpass filter (Semrock FF01-465/537/623-25) to diminish the intensity of light with wavelengths of 480 nm and longer, reducing direct excitation of the donor dye.

Fluorescence emission measurements were acquired with excitation centered at $532\,\mathrm{nm}$ or $632\,\mathrm{nm}$ (spectral width of $10\,\mathrm{nm}$). Emission spectra were collected with a spectral width of 3 nm and integration time of $200\,\mathrm{ms}$. Time-resolved measurements collected emission in a spectral band of $10\,\mathrm{nm}$ around $665\,\mathrm{nm}$ to record acceptor fluorescence or with a 3-nm band around $570\,\mathrm{nm}$ to record donor fluorescence, with an integration time of $100\,\mathrm{ms}$. The emission spectra and intensities in time traces were corrected for background from the buffer without sample.

In vitro processive single-molecule fluorescence tracking assays. In vitro assays on tetrameric motor constructs were performed as in ref. 7 with some modifications. Filaments were immobilized on the glass surface using rabbit skeletal muscle myosin (Cytoskeletal Inc, cat. no. MY02) treated with N-ethyl maleimide. Assay buffer consisted of 25 mM Tris (pH 7.5), 25 mM KCl, 1 mM EGTA, 2 mM MgCl₂ and 10 mM DTT. Blocking buffer consisted of 25 mM Tris (pH 7.5), 25 mM KCl, 1 mM EGTA, 2 mM MgCl $_2$, 10 mM DTT and 2 mg ml $^{-1}$ BSA (Sigma-Aldrich). Imaging buffer consisted of blocking buffer supplemented with an oxygen-scavenging system (0.2 mg ml⁻¹ glucose oxidase, 0.36 mg ml⁻¹ catalase and 0.4% wt/vol glucose) and an ATP regeneration system (creatine phosphokinase at 0.95 µg ml⁻¹ and 1 mM phosphocreatine) and diluted motors (final dilution 1:300 to 1:2,000 from the stock concentration of 80 nM). Unlike ref. 7, we did not supplement any of the assay buffers with 0.1% Tween. The flow protocol was as follows: (1) flow in 3.5 µl of N-ethyl-maleimide-treated muscle myosin followed by incubation for 2 min; (2) flow in 30 µl of blocking buffer followed by incubation for 2 min; (3) flow in 30 µl of diluted actin followed by incubation for 3 min; (4) flow in imaging buffer.

Channels were imaged with TIRF excitation. Fluorescence was excited with a solid-state 633 nm laser (BSR). For each channel, we first performed a quick scan across several fields of view with laser excitation at 532 nm to verify that there was a good distribution of filaments across the channel. We then performed the following image acquisition protocol on a suitable field of view: (1) turn on the red laser, acquire $\sim\!\!20$ -frame video (typical exposure time of 100 ms) of red labeled filament heads; (2) turn on the green laser, switch to the green imaging channel on Optosplit and image filaments with short video (10–20 frames, exposure of 1 s); (3) switch back to the red laser at high power (1 mW at objective back aperture) and wait for filament heads to bleach to a dim level compatible with motor imaging; (4) start image acquisition on the motor channel (red), using an exposure time of 1 s.

Motor trajectories were analyzed using the kymograph generation and analysis software KymographClear and KymographDirect¹². We manually selected only clearly polarity-labeled filaments, analyzing each of the three phases of the acquisition sequence (dark pre-, lit and dark post-) as a separate video. In KymographDirect we used the 'noise rejection' detection mode and processed results for forward and backward trajectories separately. For each run, we recorded

the average velocity of the detected trajectory, contributing one measurement to the velocity histogram. Multiple separate runs were recorded on a single motor displaying bidirectional behavior when the directional phases were picked up as separate segments by the detection algorithm.

In vitro processive single-molecule gold-nanoparticle tracking assays.

Gold nanoparticle tracking assays were performed as described in ref. ⁷, with some modifications to channel preparation and imaging conditions. Streptavidin-conjugated 50-nm gold nanoparticles (Cytodiagnostics) were washed twice in binding buffer (500 mM NaCl, 40 mM Tris pH7.5, 0.2% (vol/vol) Tween-20 (Sigma) and 1 mg ml⁻¹ BSA) by centrifugation and then resuspended by sonication. Gold nanoparticles were combined with PEG-biotin-labeled proteins at an approximately 1:1 protein:nanoparticle molar ratio and then incubated on ice for 15 min before dilution into imaging buffer. Channels were prepared as described for the in vitro processive single-molecule tracking assays with the addition of 0.2% (vol/vol) Tween-20 in the blocking buffer and imaging buffer. The polarity of Alexa 633 phalloidin-labeled actin filaments was determined using single-molecule tracking of engineered *Nicotiana tabacum* myosin XI tetramers NM11CD738_2R-1R~TET (Supplementary Figs. 1 and 2) labeled with TMR (Supplementary Video 9).

Gold nanoparticles were imaged using objective-side evanescent dark-field microscopy on a custom-built inverted microscope, with a 1.49 NA, ×60 objective (Nikon) and a 552 nm laser (Coherent OBIS 552 LS) at an irradiance of 0.8 W cm⁻² at the sample plane. Scattered light was imaged on an sCMOS camera (Andor Zyla) at 503 Hz. Actin filaments and fluorescently labeled myosins were imaged with TIRF excitation at 637 nm (Coherent OBIS 637 LX) and 552 nm, respectively, and fluorescence imaging videos were collected on an EMCCD camera (Andor iXon Ultra DU-897U). Blue illumination was provided by an LED (Thorlabs M470L2-C3) mounted above the objective. The blue LED was manually switched on to an irradiance of 17 mW cm⁻². For each channel, a video of the actin was first acquired in TIRF with excitation at 637 nm. The TMR-labeled Nicotiana tetramers were imaged in TIRF at excitation of 552 nm for directionality scoring of actin. The fluorescent tetramers were either included in the channel at the same time as the nanoparticles and imaged before imaging the nanoparticles or separately flowed in and imaged after the videos of the nanoparticles were collected. To acquire videos of the nanoparticles, the imaging was switched to dark-field scattering at excitation of 552 nm. Five to six videos of gold nanoparticles were acquired for each channel. The blue illumination was alternated between on and off for each subsequent video within a channel, remaining either on or off for the duration of each video.

Analysis of gold nanoparticle videos was performed as described in ref. 7, with some modifications. Nanoparticles were tracked using 2D Gaussian fitting with a custom routine written in C⁵⁹, and the 2D trajectory was fit with a polynomial representing the path along an actin filament. The XY positions of the nanoparticle centers were projected onto the polynomial to obtain a 1D trajectory of distance along the actin filament. Each trajectory was examined manually, and trajectories were rejected from further analysis if they displayed tracking errors (such as missed frames or tracking of nearby diffusing particles) or if the amplitude of the high-frequency noise was too large to distinguish state changes within the trajectory by eye. Steps were fit to the 1D trajectory using the Steppi change-point detection algorithm44. The following settings were used for Steppi: global unknown κ , global unknown ε , local unknown μ , $\alpha = 0$, protected state radius = 10 frames, penalty factor = 1. For step size distributions, maximum likelihood estimates of the parameters for a four-component Gaussian mixture model were obtained using the fitgmdist function in MATLAB. Standard errors for the maximum likelihood estimates were calculated using bootstrapping with 10,000 samples. Dwell time distributions were fitted to an exponential decay, omitting dwells shorter than 0.1 s, which may be undercounted due to limited experimental resolution. The standard errors of estimated decay times were calculated using the mlecov function in MATLAB.

Molecular constructs and cell lines for motor localization. Array G₁₆ was synthesized from monomeric codon-optimized ArrayG units as described in ref. 31. Using a three-piece ligation strategy, MyLOVChar4~1R~TET and ArrayG_{16X} were ligated in sequence into the multiple cloning site of a PiggyBac cumate switch system (PBQM812A-1, SBI) plasmid. This generated a MyLOVChar4~1R~TET~ArrayG_{16X} fusion plasmid, where the cumate repressor was expressed under a separate EF1 promoter as a CymR repressor-T2A-Puro cassette. To generate a binder FP plasmid, wtmGFP31 and HaloTag were ligated in sequence into a modified PiggyBac cumate switch plasmid where the cumate operator sequence was replaced with a Tet-On 3G tetracycline inducible promoter and the EF1-CymR repressor-T2A-puromycin cassette was replaced with an EF1-Tet-activator-P2A-hygromycin cassette. MyLOVChar4~1R~TET~SNAP plasmid was generated by ligating MyLOVChar4~1R~TET and a SNAP-tag in sequence into the same modified PiggyBac Tet-On 3G plasmid, except that the EF1-CymR repressor-T2A-puromycin cassette was replaced with an EF1-Tet-activator-P2A-blasticidin cassette. This was done so that, if needed, all three plasmids could be stably integrated into the same cell line using three different antibiotic resistances as selection markers. MyLOVChar4~1R~TET~mRuby3 for expression in neurons was constructed using sequence coding for the fluorescent protein mRuby 3 60 and cloned into the vector pcDNA3.1/Puro-CAG.

The transformed mouse embryonic fibroblast cell line BALB/3T3 was purchased from ATCC (CCL-163) and cultured in standard DMEM medium supplemented with 10% fetal bovine serum and penicillin–streptomycin antibiotics.

For generating stable cell lines as well as for transient transfection, cells were transfected with appropriate amounts of plasmids using the Neon Transfection system (Thermo Fisher Scientific) and the following pulse program (pulse voltage 1,350 V, pulse width 20 ms, two pulses). To generate the MyLOVChar4~1R~TET~SNAP stable cell line, BALB/3T3 cells were transfected with PiggyBAC-Tet-MyLOVChar4~1R~TET~SNAP plasmid and a Super PiggyBac transposase plasmid at 2.5: 1 ratio. Three days after transfection, the growth medium was replaced with fresh growth medium supplemented with 10 µg ml⁻¹ blasticidin. Fresh growth medium with blasticidin was added on alternate days throughout the selection process.

Molecular constructs and cell lines for controllable cargo localization.

To design an optogenetic platform for measuring directional and reversible protein-cargo delivery, we generated a two-component recruitment system. The first component, MyLOVChar4~1R~TET~SNAP~DHFR, was generated by appending a dihydrofolate reductase (DHFR) coding sequence in frame to the end of SNAP-tag from the plasmid PBTH MyLOVChar4~1R~TET~SNAP (PBTH: PiggyBac Tet-On 3G plasmid with an EF1-Tet-activator-P2A-hygromycin cassette). For the second component, either (1) the F3 subdomain of the FERM domain of mouse talin protein or (2) integrin $\beta 3$ was fused in frame to an NB113 DHFR nanobody 61 followed by a chemically inducible dimerizer sequence and eGFP, and the entire cassette was inserted at the multiple cloning site (MCS) of a PiggyBac-Tetracycline-Blasticin (PBTB) plasmid. Future applications might use the dimerizer to induce clustering using a chemical signal (TakaraBio A/C Heterodimerizer, cat. no. 635055, a rapamycin analog), but this functionality was not tested or used here.

To generate dual stable cell lines, cells from the MCF10A mammary epithelial cell line (ATCC CRL-10317) were transfected with the appropriate plasmids at a 4:1:1 Super PiggyBac transposase:component 1 plasmid:component 2 plasmid ratio using a Neon electroporator (pulse duration of 30 ms, 1,100 V, two-pulse electroporation program). Three days after transfection, the growth medium was replaced with fresh growth medium (DMEM/F12+5% horse serum supplemented with epidermal growth factor $(20\, ng\, ml^{-1})$, insulin $(10\, \mu g\, ml^{-1})$, cholera toxin $(100\, ng\, ml^{-1})$, hydrocortisone $(0.5\, \mu g\, ml^{-1})$ and penicillin–streptomycin) supplemented with $10\, \mu g\, ml^{-1}$ blasticidin and $250\, \mu g\, ml^{-1}$ hygromycin). Fresh growth medium with blastidicin and hygromycin was added on alternate days throughout the selection process.

Live-cell bulk localization assays. For bulk motor imaging in fibroblasts, the BALB/3T3 cell line expressing MyLOVChar4~1R~TET~SNAP was induced with doxycycline overnight and then live-stained with SNAP-Cell 647-SiR (New England Biolabs) according to the manufacturer's instructions. For simultaneous two-color imaging of cargo and motor, MCF10A cell lines co-expressing MyLOVChar4~1R~TET~SNAP~DHFR and NB113~integrin-β3~eGFP or NB113~FERM~eGFP were induced with doxycycline overnight and then live-stained with SNAP-Cell 647-SiR (New England Biolabs).

Cells were imaged with a Zeiss LSM 700 scanning confocal microscope. All SNAP-Cell 647-SiR labeled cells were imaged with a 639 nm laser, a 128–180- μ m pinhole, a pixel dwell time of 2.55 μ s and a ×63 plan-apochromat 1.40 NA objective.

For experiments with motors in fibroblasts, blue light stimulation was performed with either the blue 488 nm laser line of the confocal microscope at 1% of nominal power or an LED light source (Thorlabs M470L3) with a center wavelength of 470 nm. The LED was driven by a d.c. source (Thorlabs LEDD1B). For most dose–response experiments, the LED intensity was controlled by means of pulse width modulation at 500 Hz with the output of an Arduino Uno board.

For simultaneous blue light stimulation of motors and imaging of cargo in MCF10A cells, we used the 488 nm laser line of the confocal microscope at 2% laser power. Data for the four phases of the stimulus assays (pre-illumination, illumination phase 1, post-illumination and illumination phase 2) were collected in four separate videos. Three of the four phases were acquired with a typical frame rate of 14s per frame, and a lower frame rate (60 s per frame) was used for the post-illumination phase to reduce photobleaching. Video acquisition of an individual phase within a stimulus assay was started manually, ~14s after the last frame of a previous acquisition. Supplementary Table 2 provides an overview of light stimulation conditions for the complete set of experiments.

Live-cell bulk localization analysis. We quantified the accumulation of motors in puncta with the TrackMate plugin in ImageJ and subsequent analysis with custom-written scripts in Python. Tracking and puncta quantification were performed on a maximum intensity projection of the confocal *z*-stack.

To detect puncta for the experiments with motors in fibroblasts, we used the Find Maxima detector (https://imagej.net/Find_maxima_(Trackmate_module)),

with a spot diameter of 1 μ m and a noise tolerance of 100. Spots were tracked using the linear assignment problem tracker with a frame-to-frame linking distance of 2 μ m, a maximum frame gap of two frames, and maximum track segment closing distance and track segment merging distance of 4 μ m.

We quantified the population of motors in the cytoplasm by recording the mean intensity from 4–6 cells in each analyzed video. For each cell, the cytoplasmic intensity was computed as the mean value in a rectangular region of interest (ROI). The size and position of each static ROI was manually defined in ImageJ and optimized to fully remain in the cytoplasmic region of the cell for the full duration of the analysis interval. For each selected cell, we also defined a ROI in the nucleus of the cell to apply a background subtraction. Analysis was performed on a single plane chosen from the confocal *z*-stack. To prevent loss of intensity due to focal drift, we selected the plane of the *z*-stack with maximum average intensity within the set of ROIs for each frame of the video. For the dataset presented in Fig. 3f, the selection plane was constant for the duration of the experiment.

We used custom Python scripts to tabulate the tracked spots at each time point in the video and compute the average intensity of spots as a function of time, normalized to the total intensity contained in each image. For the dose–response data, we extracted the saturated value and characteristic rise time from fitting a single exponential rise to the trace (Supplementary Fig. 8).

We analyzed the accumulation of puncta in experiments on the MCF10a cells as for the experiments in fibroblasts described above, with a few modifications. We selected regions in the videos that primarily contained cells that clearly expressed both the motor and the cargo. Collections of puncta in both channels were extracted from these video crops using a Laplacian of a Gaussian (LoG) spot detection algorithm, with a spot radius of 2 pixels, a noise tolerance of 1,000 and selecting spots with a minimum contrast of 0.2. Spots were tracked using the linear assignment problem tracker with a frame-to-frame linking distance of 2 pixels, a maximum frame gap of two frames, and maximum track segment closing distance and track segment merging distance of 4 pixels. We filtered the set of found trajectories to preserve tracks of at least four frames in length, with a minimum track velocity of 1 pixel per frame; we found this empirical additional filtering useful in the epithelial cell videos to select for the puncta in protrusions, which are mobile cellular structures, excluding stuck spots. We used custom Python scripts to tabulate the tracked spots at each time point in each phase of the stimulus assay and compute the average intensity, normalized to the total intensity contained in each image. We then combined the time traces for the individual phases into a single sequence, and further normalized the traces to the value in the first frame in the trace. The resulting trajectories for each acquisition sequence (Supplementary Figs. 10 and 11) include some variable time gaps between individually acquired videos for each of the four phases of the stimulus experiment. To average the plots and place them on a common time axis in Fig. 5c,e, we used the start times of each of the two blue light stimulus phases as common reference points, and resampled the data during the post-illumination phase at common time points by interpolation before computing averages and standard deviations.

Neuron culture imaging and light stimulation. All cell culture reagents were purchased from Thermo Fisher Scientific unless otherwise specified. Glass-bottomed cell culture dishes were pre-coated with poly-D-lysine (10 µg ml-1, Sigma-Aldrich) overnight at 4°C and washed with PBS. Hippocampal neurons were dissected from embryonic day 18 rats and plated at a density of 30,000 per cm² in the plating medium (Neurobasal medium supplemented with B27, 2 mM GlutaMAX and 10% FBS). The entire medium was replaced with the growth medium (Neurobasal medium supplemented with B27, 2 mM GlutaMAX and 1% FBS) on the following day, and half of the medium was refreshed every 3-4 days. Neurons were transfected with MyLOVChar4~1R~TET~mRuby3 on the 9th day in vitro (9DIV) using Lipofectamine following the manufacturer's instructions. Cultures were maintained at 37 °C in 5% CO₂ and 100% humidity during the entire culturing period. All animal procedures were approved by the Stanford Institutional Animal Care and Use Committee. Dynamics of MyLOVChar4~1R~TET~mRuby3 were imaged in 12DIV transfected neurons using an inverted wide-field microscope (Zeiss, Axiovert 200). Before the experiment, the cell culture medium was replaced with imaging solution (Hank's balanced buffered solution supplemented with 2 mM GlutaMAX,10 mM HEPES and 1 mM D-glucose). The culture dish was equilibrated for 30 min in a 37 °C heated chamber supplied with 5% CO₂. Cells were excited with a xenon arc lamp filtered at 568/20 nm under a ×20, 0.75 NA objective lens, and images were detected with an emission filter at 620/60 nm. Timelapse images were acquired every 10 s for 80 time points. Light stimulation (~ $10\,\text{mW}\,\text{cm}^{-2}$) was applied with a blue LED lamp (Thorlabs M470L3, center wavelength of 470 nm) placed on top of the cell culture dish between time points 21 and 50 for each timelapse series.

Cell staining and imaging for single-molecule assays. For single-molecule imaging of MyLOVChar4–1R~TET~ArrayG $_{16N}$ the BALB/3T3-Tet-MyLOVChar 4~1R~TET~SNAP cell line was transiently transfected with the cumate-inducible MyLOVChar4~1R~TET~ArrayG16 and Tet-inducible mwtGFP-Halo plasmids. The cells were then induced overnight with doxycycline to ensure robust expression of MyLOVChar4~1R~TET~SNAP and the binder protein mwtGFP-Halo. Cumate was not added, as the leaky expression of

MyLOVChar4~1R~TET~ArrayG $_{16X}$ under zero cumate induction was ideal for single-molecule imaging. Overexpression of MyLOVChar4~1R~TET~SNAP ensured that the MyLOVChar4~1R~TET~ArrayG $_{16X}$ were tetramers. Single molecules were imaged in HILO using a TIRFM set-up on an Olympus CellTIRF system through a 1.49 NA ×100 objective and ×1.6 Optovar magnifier. For imaging MyLOVChar4~1R~TET~ArrayG $_{16X}$ /mwtGFP-Halo we used a BrightLine 482/18 excitation filter and a BrightLine 525/45 emission filter. Images were acquired at 10 Hz on an Andor iXon Plus EMCCD camera.

Live-cell single-molecule analysis. We tracked single particles using the MATLAB software package μ -track, implementing tracking as a linear assignment problem algorithm 2. For detection, we used an initial Gaussian standard deviation of 2 pixels, rolling window averaging of three frames, iterative estimation of Gaussian standard deviation and iterative Gaussian mixture modeling with residuals, distance and amplitude parameters set to 0.05. For tracking, we used a problem dimensionality of 2, a maximum gap to close of four frames, minimum length of track from the first step of one frame, and the options for segment splitting and segment merging turned on. We used Brownian and directed motion models as the cost functions for frame-to-frame linking, gap closing, merging and splitting, and a Kalman filter function. For initial motion analysis in μ -track, we used a problem dimensionality of 2 and an α value of 0.1 to analyze asymmetric tracks.

Extracted tracks were then analyzed with custom-written MATLAB scripts to extract the parameters of all tracks classified as directed motion, with a minimum length of 20 frames. Within a track, we designated mobile and stalled time points by simple thresholding. A histogram of velocities of the mobile phases was populated with the average velocities of all contiguous mobile sections of the tracks that were longer than four frames, recording the average velocity of each section as one data point.

Replication and reproducibility. All in vitro assay data were collected on two or more independent preparations of the assays. Additionally, for direct comparisons of properties between different molecular motor constructs (Fig. 1f), data were collected using at least two independent protein preparations for each construct. All data in live cells were collected over at least two independent biological replicates, on separate days.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this Article.

Data availability

Data for all results reported in this study are freely available as follows. The cryo-EM map and atomic model for the myosin XI–F-actin complex have been deposited under accession numbers EMDB-22808 and PDB 7KCH, respectively. Source data are available on the journal website as spreadsheet files linked to Figs. 1–5 and Extended Data Figs. 3 and 4. Source data for Supplementary Figs. 3–11 are available from the Stanford Digital Repository (https://doi.org/10.25740/65j8-6114). All unique biological materials generated for this study, including plasmids for purification of engineered motors and for live-cell experiments, are available from the authors on request. Source data are provided with this paper.

Code availability

Custom code used in this study is freely available from the Stanford Digital Repository (https://doi.org/10.25740/65j8-6114).

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Author contributions

P.V.R., M.N., R.C. and V.T.V. designed and produced engineered myosin constructs. P.V.R., S.Z., M.N. and V.T.V. designed and performed in vitro measurements. P.V.R. and S.Z. developed analysis code and analyzed in vitro data. R.P.G. designed and performed research in fibroblasts and epithelial cells, including the design and production of constructs and cell lines and consulting on analysis approaches. P.V.R. analyzed live-cell data and consulted on live-cell imaging. R.G. designed and performed cryo-EM research and analyzed structural data. L.N. designed and performed research in neurons. R.R.E., N.A. and A.E.C. designed and purified the FRET construct and contributed structural modeling expertise. Z.B., J.T.L., G.M.A., M.Z.L. and P.-S.H. supervised research. All authors discussed the results and contributed to writing the manuscript.

Competing interests

The authors declare no competing interests.

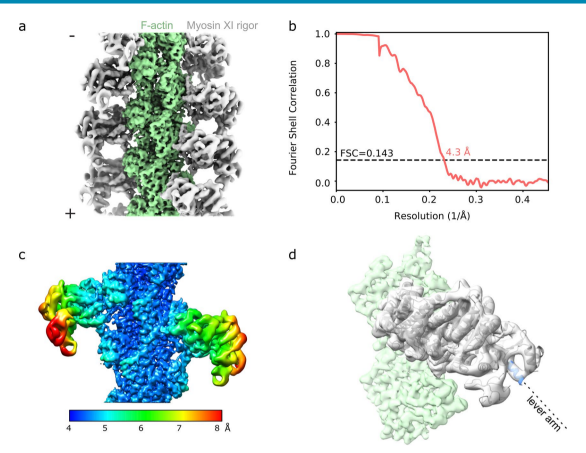
Additional information

Extended data is available for this paper at https://doi.org/10.1038/s41589-021-00740-7.

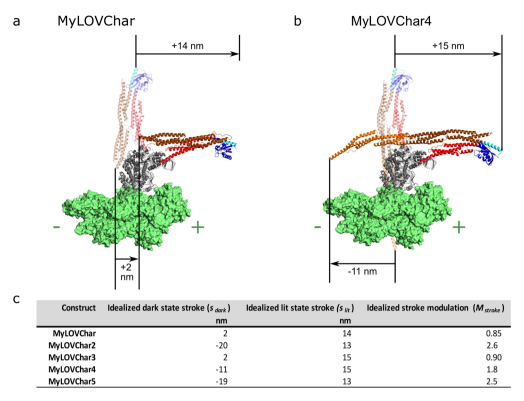
 $\label{thm:contains} \textbf{Supplementary information} \ The online version contains supplementary material available at $$https://doi.org/10.1038/s41589-021-00740-7.$

Correspondence and requests for materials should be addressed to Z.B.

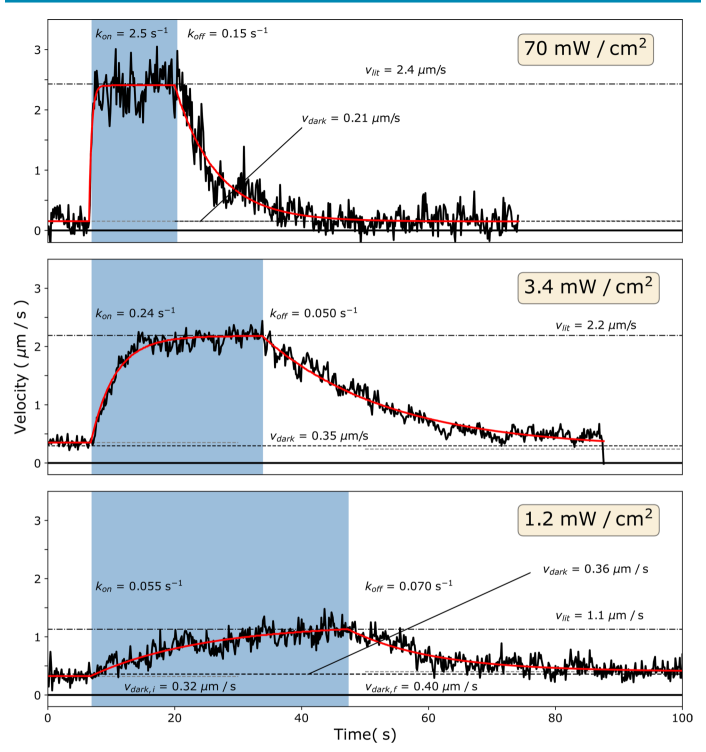
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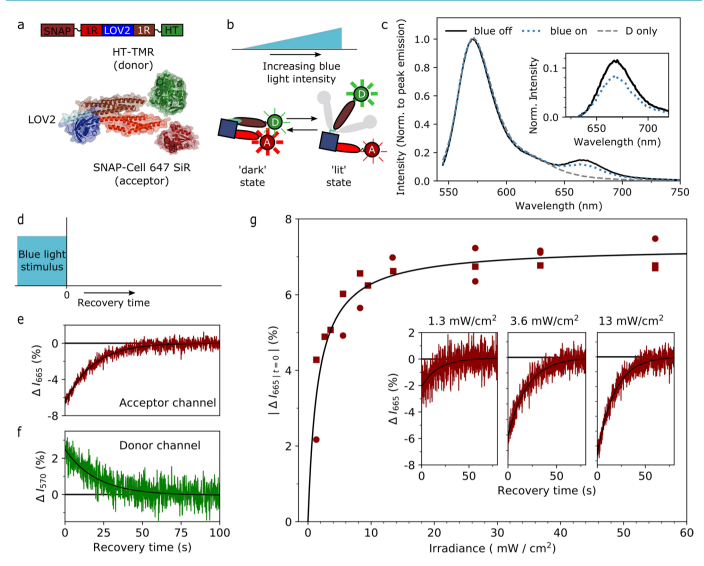
Extended Data Fig. 1 | Cryo-EM analysis of F-actin decorated with rigor state myosin XI. a, Cryo-EM map of the rigor state of the *Chara* myosin XI catalytic domain $CM11_{L2+4}$ CD_{746} bound to F-actin, reconstructed at 4.3 Å. **b**, The gold-standard Fourier shell correlation (FSC) curve for the 3D reconstruction. The average resolution is estimated at the FSC value of 0.143. **c**, Local resolution estimation of the 3D reconstruction. **d**, Cryo-EM map of two actin and one myosin subunits with the ribbon model of myosin fitted. The terminal helix of the myosin converter domain (residues 738-746) is highlighted in cornflower blue; this helix can be extended by a fused lever arm as indicated by the dashed line.



Extended Data Fig. 2 | Structural models and predicted stroke vectors. a,b, Structural models of MyLOVChar and MyLOVChar 4, displaying representative pre-stoke (transparent) and post-stroke (solid) models from a set of 50 top-scoring structures modeled using RosettaRemodel (Methods and Supplementary Table 1), with annotated actin polarity. Post-stroke models are based on an actin-bound rigor stroke structure of *Chara corallina* myosin XI structure (Fig. 1b and Extended Data Fig. 1). Pre-stroke structure models built from myosin V pre-stroke crystal structure were aligned to the actin-bound myosin XI rigor. Idealized power stroke sizes and directions in the dark state (arrows below structures) and in the lit state (arrows above structures) were calculated by projecting the motion of the effective tip of the lever arm along the actin filament for the 50 top-scoring structures, and recording the mean. The effective tip of the lever arm was taken as the position of the C-terminal residues in the dark state, and as the position of the last rigid element on the LOV2 domain in the lite state (G516) **c**, Tabulation of calculated stroke sizes and optical stroke modulation M_{stroke} for structural models of MyLOVChar5. Negative values indicate a stroke directed to the actin minus end. The stroke modulation M_{stroke} was calculated as 1- $s_{\text{dorh}}/s_{\text{lit}}$. A modulation larger than 1 indicates a directional reversal of the stroke.



Extended Data Fig. 3 | **Velocity traces of gliding filament assays on MyLOVChar4 at variable blue light intensity.** The traces are selected from the dataset used to compose Fig. 1h in the main text. Each trace displays the frame-to-frame velocity at the annotated blue light irradiance (box in the top right corner), averaged over all tracked polarity-labeled filaments in 2-3 consecutive repeats of the stimulation sequence. The initial dark velocity $v_{dark,i}$ was taken as the mean velocity of the time before the start of the blue light stimulus, indicated by the left gray dashed lines, and annotated for the bottom plot. The subsequent parts of the trace were globally fit to an exponential rise and decay, extracting the characteristic rates k_{on} and k_{off} , the steady-state lit velocity $v_{dark,f}$ (black dash-dotted lines), and the final velocity $v_{dark,f}$ (right gray dashed lines, annotated for bottom plot). In Fig. 1e and Fig. 1h the dark velocity v_{dark} (black dashed lines) was reported, taken as the mean of $v_{dark,f}$ and $v_{dark,f}$.



Extended Data Fig. 4 | Dose-dependent conformational changes in a minimal LOV2-containing lever arm construct. a, Block diagram and idealized molecular cartoon of a minimal lever arm construct with N-terminal SNAP-tag and C-terminal HaloTag fusions (approximate positions shown). HaloTag was labeled with TMR (donor) and SNAP-tag with SNAP-Cell 647-SiR (acceptor) for fluorescence measurements. b, Schematic of light-dose-dependent population of conformational states of the construct. Transparent elements in the lit state indicate increased flexibility. Positions of the fluorescent dyes are annotated (D: donor, A: acceptor). c, Fluorescence emission spectra normalized to the peak donor emission. Inset: extracted acceptor emission, obtained by subtracting the normalized emission from a donor-only sample from the normalized emission of the FRET sample. d, Schematic of the light stimulus experiment in bulk in solution. e-f,Time traces of fluorescence intensity immediately after a removal of a blue light stimulus (irradiance: 10 mW/cm²), with continued excitation of the donor dye. e, Fluorescence intensity collected at the emission maximum of the acceptor (665 nm). f, Fluorescence intensity collected at the emission maximum of the donor (570 nm), recorded on an independent run of the stimulus sequence, on the same sample as d using the same blue light irradiance. g, Amplitude of the normalized intensity difference of acceptor channel after recovery from a light stimulus as function of irradiance of the blue light, showing two replicate datasets (circular and square data points) taken on different days. Insets: time-dependent traces for measurements at the annotated irradiances. Black lines in e-f and insets in g are single exponential decays with amplitude A and characteristic time t. Listed in pairs (A, t), parameter values are e: (-7 %, 18 s), f: (+3 %, 20 s), g (left): (-2 %, 20 s), g (center): (-5 %, 20 s), g (right): (-7 %, 18 s) s). Black line in g is a saturation curve with A=7 % at saturation, with half-saturat

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Reporting Summary

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For	all st	atistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.
n/a	Cor	nfirmed
	\boxtimes	The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement
	\boxtimes	A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
\boxtimes		The statistical test(s) used AND whether they are one- or two-sided Only common tests should be described solely by name; describe more complex techniques in the Methods section.
\boxtimes		A description of all covariates tested
X		A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons
	\boxtimes	A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)
\boxtimes		For null hypothesis testing, the test statistic (e.g. <i>F</i> , <i>t</i> , <i>r</i>) with confidence intervals, effect sizes, degrees of freedom and <i>P</i> value noted <i>Give P values as exact values whenever suitable.</i>
\boxtimes		For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
\boxtimes		For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
\boxtimes		Estimates of effect sizes (e.g. Cohen's d, Pearson's r), indicating how they were calculated
	'	Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection

-cryo-electron microscopy (image stack recording): Leginon software (version 3.2).

-in vitro microscopy (gliding motility and single-molecule tracking): custom Labview program (Labview 8.5) for data acquisition on a Nikon TIRF microscope.

-in vitro microscopy (gold nanoparticle tracking): custom Matlab program (Matlab 2016a) for data acquisition on a custom modified Nikon microscope

-live cell microscopy: Zeiss Zen software (2012 SP5 FP1) to perform data acquisition on a scanning confocal microscope (LSM 700 scanning confocal microscope), and a custom Arduino macro to operate a blue LED. Live cell single-molecule tracking experiments were performed on an Olympus CellTIRF system with custom acquisition code developed in the Liphardt lab.

Data analysis

Software is listed below; see Methods for details of how software was used. Custom code used in this study is freely available from the Stanford Digital Repository (https://doi.org/10.25740/65j8-6114).

-cryo-electron image analysis: RELION 3.0 image processing pipeline; MotionCor2 (version 1.1.0) for motion correction; CTFFIND4 (version 4.1) for CTF estimation; UCSF Chimera (version 1.14) and ChimeraX (version 0.9);

-cryo-electron model building: I-TASSER (https://zhanglab.ccmb.med.umich.edu/I-TASSER), UCSF Chimera (version 1.14); Rosetta (version 3.10); coot (version 0.9.1); Phenix.real_space.refine in Phenix suite (version 1.18.2-3874)

-MyLOVChar1-5 modeling: Rosetta (version 3.8), PyMOL (version 2.2.3)

-gliding motility analysis: custom Matlab scripts (Matlab 2018a); ImageJ (version 2.0.0-rc-69/1.52n) with plugins IJ-ridge-detect (version 1.4.0, by Wagner, Hiner and Raynaud) and Trackmate (version 4.0) and custom Jython scripts; custom software written in Python (Python 3.7) modified from Tural Aksel's FAST software package (in Python 2.7).

-in vitro processive single-molecule tracking analysis: KymographClear (version 2.0) and KymographDirect (version 2.1a for Mac OS, a custom version provided by Pierre Mangeol).

-live cell localization analysis: ImageJ with custom Jython scripts and TrackMate (version 4.0) using the TrackMate module FindMaxima (version 1.0.0); custom Python scripts.

-live cell single-molecule tracking: micro-track (version 2.2, running in Matlab 2018a) software from Danuser lab (UT Southwestern); custom software in Matlab and Python.

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors/reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.

Data

Policy information about availability of data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

Data for all results reported in this study are freely available as follows. The cryo-EM map and atomic model for the myosin XI-F-actin complex have been deposited in the EMDB (22808) and PDB (7KCH) respectively. Source data are available on the journal website as spreadsheet files linked to figures 1-5 and Extended Data figures 3-4. Source data for supplementary figures 3-11 are available from the Stanford Digital Repository (https://doi.org/10.25740/65j8-6114).

Field-spe	ecific reporting
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Life sciences	Behavioural & social sciences Ecological, evolutionary & environmental sciences
For a reference copy of t	the document with all sections, see <u>nature.com/documents/nr-reporting-summary-flat.pdf</u>
Life scier	nces study design
All studies must dis	sclose on these points even when the disclosure is negative.
Sample size	No predetermined sample sizes were used; sample sizes for various assays are reported in the manuscript, with distributions of data presented to support claims, and repetitions performed to establish reproducibility. Sample sizes were chosen to be sufficient for clear assessment of molecular behaviors and experimental variability.
Data exclusions	Data exclusions are described in the Methods, including movies in which filament density precluded automated tracking, exclusion of stuck filaments according to defined criteria, and thresholds applied in single-molecule analysis; criteria were empirical and not predetermined.
Danliastias	All reported approximants were replicated at least twice. Additionally, in vitra gliding motility assays on anticolly controlled a mater proteins.
Replication	All reported experiments were replicated at least twice. Additionally, in vitro gliding motility assays on optically controllable motor proteins were performed with at least two independent preparations of purified motor proteins, over at least two individual measurement days. All experiments in cells were replicated on at least two separate days, and include results from multiple cells as described in the manuscript. All attempts at replication were successful, and are included as data in the manuscript.
Randomization	Our study does not involve allocation into different experimental groups.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

conditions. We sought to avoid bias and subjectivity by consistent application of quantitative analysis to extract parameters.

We did not use blinding in our study. Blinding was not necessary due the largely automated data pipeline used to compare samples and

Materials & experimental sys	stems Me	Methods	
n/a Involved in the study	n/a	Involved in the study	
Antibodies	\bowtie	ChIP-seq	
Eukaryotic cell lines	\boxtimes	Flow cytometry	
Palaeontology	\bowtie	MRI-based neuroimaging	
Animals and other organisms			
Human research participants			
Clinical data			
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Antibodies

Blinding

Antibodies used

-anti-FLAG resin (Sigma-Aldrich A2220)

-anti-GFP (Millipore Sigma MAB3580)

Validation

Antibodies were used only for protein purification (anti-FLAG) and for surface attachment of motors in gliding filament assays (anti-GFP), following established methods using the same antibodies in prior work including Nakamura et al. (Nature Nanotechnology 2014). The manufacturer states that the anti-FLAG resin has been used "..for the purification of FLAG fusion proteins." The manufacturer states the anti-GFP has been "validated for use in ELISA, IC, IH, & WB" and reports that "MAB3580 reacts with Green Fluorescent Protein (GFP). The antibody reacts with various forms of GFP including eGFP including those found in vectors supplied by Clontech and Invitrogen. There is no cross-reactivity to RFP. YFP has not been tested. This antibody works well for immunohistochemistry."

Eukaryotic cell lines

Policy information about cell lines

Cell line source(s)

We purchased mouse embryonic fibroblast cell line BALB/3T3 and MCF10A mammary epithelial cell line (CRL-10317) from ATCC, as described in the Methods.

Authentication

The cell lines were not authenticated in our lab.

Mycoplasma contamination

Dual stable MCF10A cell lines tested negative for mycoplasma contamination. Fibroblasts were not tested.

Commonly misidentified lines (See ICLAC register)

No commonly misidentified cell lines were used.

Animals and other organisms

Policy information about studies involving animals; ARRIVE guidelines recommended for reporting animal research

Laboratory animals

Species: Rat, Domestic. Strains: Sprague Dawley. Sex: both male and female. Age: embryonic day 18.

Wild animals

Not applicable.

Field-collected samples

Not applicable.

Ethics oversight

All animal procedures were approved by the Stanford Institutional Animal Care and Use Committee.

Note that full information on the approval of the study protocol must also be provided in the manuscript.