Kinesin-1–powered microtubule sliding initiates axonal regeneration in *Drosophila* cultured neurons

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**ABSTRACT** Understanding the mechanism underlying axon regeneration is of great practical importance for developing therapeutic treatment for traumatic brain and spinal cord injuries. Dramatic cytoskeleton reorganization occurs at the injury site, and microtubules have been implicated in the regeneration process. Previously we demonstrated that microtubule sliding by conventional kinesin (kinesin-1) is required for initiation of neurite outgrowth in *Drosophila* embryonic neurons and that sliding is developmentally down-regulated when neurite outgrowth is completed. Here we report that mechanical axotomy of *Drosophila* neurons in culture triggers axonal regeneration and regrowth. Regenerating neurons contain actively sliding microtubules; this sliding, like sliding during initial neurite outgrowth, is driven by kinesin-1 and is required for axonal regeneration. The injury induces a fast spike of calcium, depolymerization of microtubules near the injury site, and subsequent formation of local new microtubule arrays with mixed polarity. These events are required for reactivation of microtubule sliding at the initial stages of regeneration. Furthermore, the c-Jun N-terminal kinase pathway promotes regeneration by enhancing microtubule sliding in injured mature neurons.

**INTRODUCTION** Neurons are highly polarized cells with long axons extending far away from the cell body. Their length makes them highly susceptible to mechanical damage, and it is crucial for an injured neuron to regenerate its axon to recover its function. Previous work on axon regeneration in *Drosophila* neurons in vivo revealed that molecular mechanisms underlying axonal regeneration include signaling pathways and cytoskeletal reorganization (Stone et al., 2010, 2012; Xiong et al., 2010, 2012; Xiong and Collins, 2012; Chen et al., 2012; Ghannad-Rezaie et al., 2012; Song et al., 2012; Klinedinst et al., 2013). However, in vivo analysis does not allow for high-resolution imaging of the regeneration process and limits application of pharmacological inhibitors for the analysis of regeneration mechanisms. In this study, we developed an in vitro axotomy technique in cultured *Drosophila* neurons and manipulated the regeneration process using both *Drosophila* genetics and pharmacological drugs.

There are two occasions when major neurite outgrowth occurs: during neurogenesis from a spherical progenitor cell to a differentiated neuron, and during neuronal regeneration after injury. Previously, we demonstrated that kinesin-1–powered microtubule sliding drives initial neurite outgrowth in young neurons by pushing the peripheral membrane forward and that this sliding is developmentally down-regulated in mature neurons once neurite outgrowth is completed (Lu et al., 2013b). The in vitro axotomy technique allowed us to show that axotomy induces a fast spike of calcium, which results in local depolymerization of microtubules. This depolymerization is followed by formation of new microtubule arrays of mixed polarity, leading to reactivation of kinesin-1–driven microtubule sliding and initiation of regeneration. Furthermore, we observed that sliding reactivation is modulated by c-Jun N-terminal kinase (JNK) pathway, a known stress-activated protein kinase cascade.
RESULTS

Kinesin-1–powered microtubule sliding drives neurite outgrowth in injured motor neurons from the larval brain

As we previously described, kinesin-1–powered microtubule sliding drives initial neurite outgrowth in young growing neurons, but sliding mostly stops in mature neurons (Lu et al., 2013b). We hypothesized that the same mechanism could be reactivated during neuronal injury, providing mechanical force for initial regeneration. We first performed an experiment to assay the role of microtubule sliding in neurons after massive damage. In Drosophila third-instar larva, motor neurons extend their axons from the ventral ganglion to target muscles to form neuronal muscular junctions that control muscle movement (Figure 1, A and B; Keshishian et al., 1996; Pilling et al., 2006). We dissociated motor neurons from the dissected larval brains (Egger et al., 2013). These motor neurons lose their neurites during dissection and dissociation (Figure 1, C and D) and therefore represent a good model of massive neuronal injury. Within 2–3 d in culture, these dissociated motor neurons regrow long (>50 μm) neurites (Figure 1E). In these long neurites, we observed active organelle transport as shown by a mitochondrial marker (upstream activation sequence [UAS]-Mito–green fluorescent protein [GFP]) under a motor neuron–specific promoter, D42-Gal4 (Pilling et al., 2006; Supplemental Figure S1, A and A′). We expressed an axonal marker, enhanced GFP-tagged synaptotagmin (Syt-eGFP; Zhang et al., 2002), in motor neurons and found that Syt-positive vesicles usually accumulated in the longest neurite (Supplemental Figure S1B). We also expressed a dendritic marker, DenMark (Nicolai et al., 2010), and found that it was excluded from the longest neurite (Supplemental Figure S1C). Furthermore, we examined the microtubule polarity using an EB1-GFP construct under D42 driver and showed that the longest neurites invariably contain microtubules of uniform polarity (plus-ends out; Supplemental Figure S1, D and D′), consistent with known polarity of microtubules in axons of motor neurons in vivo (Stone et al., 2008). Taking the results together, we conclude that motor neurons dissociated from the larval brain are capable of regenerating axons (the longest neurite) in culture.

To examine microtubule movement in motor neurons after dissociation, we expressed the photoconvertible EOS-tagged α-tubulin (Lu et al., 2013b) under D42 driver and converted it in the cell body from green to red, thus applying fiduciary marks on microtubules. By following photoconverted microtubule segments in the red channel, we observed a very high level of microtubule sliding in these freshly plated (2–4 h) neurons (N = 29). Microtubules from the photoconverted areas penetrated the neurites and underwent extensive bending, looping, and sprouting to growing neurites (Figure 2, A and B, and Supplemental Video S1). However, after the regeneration is completed, microtubule sliding becomes undetectable both in the cell body and in neurites (Figure 2, C and D; N = 29; Supplemental Video S2).

As seen during initial neurogenesis, microtubule sliding is completely dependent on kinesin-1 heavy chain (KHC; Lu et al., 2013b). Maternal load of wild-type KHC mRNA from KHC heterozygous mutant mothers allows KHC homozygous mutant flies to complete most of the embryonic and larval neurogenesis, until second-instar larvae, when maternaly loaded mRNA gets completely depleted (Saxton et al., 1991; Hurd and Saxton, 1996; Brendza et al., 2000). Therefore we cultured larval brain neurons from KHC mutant third-instar larvae with the maternal load of KHC already depleted to address the role of KHC in regeneration. KHC mutant larval brain neurons have very little microtubule motility (Figure 3, A–A′, and Supplemental Video S3; N = 20), and all failed to regenerate axons after dissociation and plating (Figure 3, B, C, C′, and E).

Kinesin-1 in neurons is required for both organelle transport and microtubule sliding (Barlan et al., 2013; Lu et al., 2013b); thus inhibition of regeneration in KHC mutant larval brain neurons could potentially be caused by disruption of organelle transport. Therefore we also tested the role of organelle transport during regeneration. Because kinesin-1 and cytoplasmic dynein are interdependent for organelle transport (Hancock, 2014), inhibition of cytoplasmic dynein abolishes kinesin-1–mediated transport of multiple cargoes, including synaptotagmin-containing synaptic vesicles (Martin et al., 1999), mitochondria (Pilling et al., 2006), peroxisomes (Kural et al., 2005; Ally et al., 2009), neurofilaments (Uchida et al., 2009), lipid droplets (Gross et al., 2002; Shubeita et al., 2008), and ribonucleotide granules (Ling et al., 2004). We expressed RNA interference (RNAi) against the Drosophila dynein heavy chain (DHC64C-RNAi) in the larval nervous system, and this RNAi led to larval posterior paralysis and complete pupal lethality, which is similar to the phenotypes of Dhc64C transheterozygotes combining a hypomorphic allele Dhc64C<sup>−</sup>−10 with either a Dhc64C deletion Df(3L)10H or an amorphic allele Dhc64C<sup>−</sup>−19 (Martin et al., 1999). Cultured DHC64C-RNAi larval brain neurons display no major defects in axonal outgrowth compared with control (Figure 3, B, D, and E). However, the kinesin-1–dependent mitochondrial movement in these DHC64C-RNAi larval brain neurons is significantly diminished compared with the control (Supplemental Figure S2, A–A′ and B–B′). Taking the results together, we conclude that the failure of regeneration in KHC mutant neurons does not result from inhibition of organelle transport, because DHC-RNAi efficiently blocks organelle transport but not axonal regeneration in larval brain neurons. These data strongly suggest that kinesin-1–powered microtubule sliding drives initial axon regeneration after traumatic axonal injury.

Microtubule sliding is activated after axotomy

To study axonal local real-time response to injury, we performed mechanical axotomy in individual neurons. We cultured neurons from dissociated gastrulation embryos (Lu et al., 2013a,b) and...
there is no visible microtubule sliding in intact mature neurons (4 d in culture; Supplemental Figure S3, D and D'; Lu et al., 2013b), robust sliding is reacti-
vated after injury: the microtubule segments slide into the regenerating tips, and sliding reactivation consistently correlates with the
initiation of axon outgrowth after injury (Figure 4, C–D', orange arrowheads, and Supplemental Video S5). In addition to mi-
rotubules pushing against the tips of the growing processes, we also observed micro-
tubules moving in the retrograde direction, implying bidirectional sliding of microtu-
bules (Figure 4, C' and D', blue arrowheads, and Supplemental Video S5). Taken to-
gether, these results support the idea that microtubule sliding is reactivated by injury to initiate axonal regrowth.

Axotomy leads to a fast Ca^{2+} influx and formation of microtubules of mixed polarity
We next investigated what the initial injury signal could be from the injury site that
causes the axon to regenerate. Calcium has been implicated as an essential player in the
neuronal injury process (Ziv and Spira, 1995, 1997; Gitler and Spira, 1998; Kamber et al.,
2009; Ghosh-Roy et al., 2010). We used a calcium reporter, GCaMP3 (Tian et al.,
2009), to monitor the intracellular calcium level upon injury. We observed a rapid cal-
cium influx initiating at the injury site that
spreads along the neurite (Figure 5, A and A', and Supplemental Figure S4A) and
reaches the cell body within seconds (Figure 5A'', Supplemental Figure S4A, and Supple-
mental Video S6). The calcium signal usually drops back to the normal level within 1–2
min (Figure 5A''').

Calcium has been shown to depolymer-
ize microtubules (Marcum et al., 1978; Keith
et al., 1983; O’ Brien et al., 1997). Therefore
we wanted to determine whether microtubules are depolymerized
by the calcium influx after injury. We imaged elav>tdEOS-\(\alpha\)-tub neu-
rons in green channel during the axotomy procedure and found very
fast microtubule disassembly in the distal axon near the injury site
(Figure 5, B and B', and Supplemental Video S7) without any signifi-
cant axon retraction (Figure 5B''), whereas much less depolymerization
was detected in the cell body or the proximal axon (Figure 5, B,
B', and C). We further found that the microtubules at the injury site
recovered within 2 h after the fast depolymerization (Supplemental
Figure S5A).

We then examined whether this calcium influx is necessary for
initiation of axonal regeneration. We added 10 mM ethylene glycol
tetraacetic acid (EGTA) to the medium to chelate extracellular cal-
cium and, using elav>GCaMP3, demonstrated that this concentra-
tion of EGTA completely blocks calcium influx after axotomy
(Supplemental Figure S4B; compare to Supplemental Figure S4A;
Supplemental Video S6). We further examined microtubules upon
axonotomy in the presence of 10 mM EGTA and found that without
the calcium influx, microtubules stay intact after injury (Figure 5C,
Supplemental Figure S4, C–C''', and Supplemental Video S7).

**FIGURE 2:** Microtubule sliding promotes neurite outgrowth during larval motor neuron
geneneration. (A, B) Motor neurons expressing photoconvertible tdEOS-tagged \(\alpha\)-tubulin 84B
under the motor neuron–specific D42 driver (D42-tdEOS-\(\alpha\)-tub84B) at 2–4 h in culture after
plating, showing a high level of microtubule sliding. (C, D) Motor neurons (D42-tdEOS-\(\alpha\)-tub84B)
at >48 h in culture after plating, showing no detectable microtubule sliding in the cell body
(C) or in the neurite (D). Column 1 shows the green signal of tdEOS-\(\alpha\)-tub84B before
photoconversion, with the photoconversion zone marked by a purple circle; columns 2–4 show
the red signal of tdEOS-\(\alpha\)-tub84B after photoconversion in the indicated areas (purple circles).
Note that in A and B, photoconverted microtubules move outside the photoconversion area.
Scale bar, 5 μm.

removed the distal portions of axons by using a glass needle oper-
ated by a micromanipulator (see details in Materials and Methods;
Figure 4A). We first imaged microtubules using an endogenous
GFP-tagged microtubule-associated protein, Jupiter-GFP (Morin
et al., 2001; Karpova et al., 2006; Lu et al., 2013b), and found that
there is little neurite outgrowth or microtubule motility in intact 4-d-
old mature neurons (Supplemental Figure S3, A–C). In contrast, af-
after axotomy, regenerating axons contain extensively looping micro-
tubules (Figure 4, B and B', and Supplemental Video S4), a strong
indication of microtubule sliding reactivation. Furthermore, we
noticed that these microtubules pushed against the very tip of the
neurite membrane during the regeneration process (compare phase
images in Figure 4B to Jupiter-GFP images in Figure 4B', Supple-
mental Video S4).

To visualize directly microtubule sliding, we examined photocon-
verted microtubule segments in neurons expressing photoconvert-
ible tdEOS-\(\alpha\)-tub84B under a panneuronal-specific driver, elav
(Campos et al., 1987; Robinow and White, 1988), using total internal
reflection fluorescence (TIRF) microscopy. We found that although
there is no visible microtubule sliding in intact mature neurons (4 d
Further, no regeneration can be detected after axotomy in the presence of 10 mM EGTA (Figure 6A), indicating that calcium influx after injury is necessary to initiate regeneration.

One of the possible roles of calcium during regeneration could be depolymerization and subsequent reorganization of the microtubule network. To observe directly the reorganization of the microtubule network, we used a plus-end marker, 

ubiquitin-Jupiter-mCherry. (B) A control neuron (Khc27/Khc23) neuron (-72 h after plating) has very short neurites. (D) A DHC64C-RNAi mutant neuron (-72 h after plating) has a long axon. (E) Quantification of the axon length in control, Khc mutant (Khc27/Khc23), and DHC64C-RNAi (-72 h after plating). The average (\( \pm 95\% \) confidence interval [CI]) for the control is 86.8 ± 5.5 μm (n = 56); the average (\( \pm 95\% \) CI) for the Khc mutant is 14.7 ± 2.2 μm (n = 80); the average (\( \pm 95\% \) CI) for DHC64C-RNAi is 88.5 ± 5.6 μm (n = 76). Unpaired t test between control and Khc mutant neurons: ***p < 0.0001; unpaired t test between Khc mutant and DHC64C-RNAi neurons: ***p < 0.0001; unpaired t test between control and DHC64C-RNAi neurons: p = 0.6693 (nonsignificant). Scale bar, 5 μm.

>1 h after injury induction (Supplemental Figure S5B). This observation is in agreement with previous studies showing microtubule polarity change upon injury (Kamber et al., 2009; Song et al., 2012). We propose that microtubule repolymerization after Ca\(^{2+}\)-induced disassembly occurs on the residual nucleation sites without directional bias and results in the formation of new mixed-polarity microtubule arrays.

To analyze the role of microtubule depolymerization and the subsequent polarity change in the early stages of axonal regeneration, we performed axotomy in the presence of a low concentration of vinblastine (10 nM). This substoichiometric amount of the drug is sufficient to block microtubule dynamics (Supplemental Figure S6, A and B; Lu et al., 2013b) but does not cause depolymerization of the existing microtubule network (Supplemental Figure S6, C and D). We observed that inhibition of microtubule dynamics by vinblastine completely prevents regeneration (Figure 6A and Supplemental Video S8). There are two possible explanations for the inhibition by vinblastine: 1) tubulin polymerization is the major force driving initial regeneration, and 2) microtubule polarity change is essential for regeneration initiation. We reasoned that since the microtubule polarity change occurs promptly after axotomy, inhibition of microtubule dynamics only at the earlier stage or only at the later stage by vinblastine would allow us to distinguish between these two possibilities. Therefore first we performed the axotomy in presence of 10 nM vinblastine and waited 2 h before we washed out the drug. This vinblastine treatment eliminated formation of mixed-polarity arrays near the injury site (Supplemental Figure S6E) and abolished regeneration (Figure 6A). Second, we added 10 nM vinblastine 3–4 h after the axotomy procedure, when formation of mixed-polarity arrays is already completed. Unlike addition of vinblastine before the time of injury, this delayed addition did not block microtubule sliding or regeneration (Figure 6A); instead, we observed microtubule looping and sprouting in the absence of microtubule dynamics (Figure 6, D–D′′′, and Supplemental Video S8), which provides strong evidence of microtubule sliding after axotomy. Therefore we conclude that mixed-polarity microtubules are essential for the initiation of axonal regeneration.

Kinesin heavy chain drives sliding of mixed-polarity microtubules

We next examined whether KHC is responsible for microtubule sliding after injury, as it is during initial neurite outgrowth (Lu et al., 2013b). We specifically knocked down KHC in the Drosophila nervous system by expressing RNAi against KHC (see Materials and Methods). Knockdown of KHC leads to larval growth defects, locomotion abnormality, and eventually 100% lethality before reaching...
brevin D treatment abolished kinesin-1–dependent mitochondrial movement (Supplemental Figure S6, F and G; Firestone et al., 2012; Lu et al., 2013b; Sainath and Gallo, 2014), as well as kinesin-1–dependent peroxisome transport (Firestone et al., 2012). However, ciliobrevin D does not affect kinesin-1–mediated microtubule sliding (Lu et al., 2013b). Therefore ciliobrevin D treatment allows us to distinguish between two roles of kinesin-1. In ciliobrevin D–treated neurons, we consistently observed regeneration events at a level comparable to control (Figure 6A). These data exclude the possibility that the failure of regeneration in KHC-RNAi neurons is caused by inhibition of organelle transport. Furthermore, these data indicate that kinesin-1, not cytoplasmic dynein, is the main driver of microtubule sliding during initial regeneration after axotomy.

JNK activation stimulates microtubule sliding and promotes regeneration

We showed that change of microtubule polarity caused by calcium influx is necessary for kinesin-1–powered microtubule sliding during initial regeneration. However, microtubule reorganization alone is probably not sufficient for sliding reactivation, since microtubule polarity changes occur within 30 min of axotomy, but we usually observe initial regeneration only after 2 h in culture. Therefore we further investigated the role of an injury-response signaling
FIGURE 5: Axotomy leads to fast Ca$^{2+}$ influx and microtubule depolymerization. (A–A″′) Signal heat maps of a neuron-specific calcium reporter (elav-GCaMP3) before and after axotomy, showing that a fast spike of calcium occurs upon mechanical axotomy. Axotomy is indicated by the white arrowheads (A′−A″′). (B, B′) Fast microtubule depolymerization occurs upon axonal injury in an elav-tdEOS-αtub84B neuron (green channel). (B″) Postcutting phase image of B′. (C) Quantification of relative fluorescence intensity of green tdEOS-αtub84B signal at two sites of the axon, near the cutting site and near the cell body, in control neurons (red and blue boxes in B and B′, respectively) and in 10 mM EGTA-treated neurons (purple and green boxes in Supplemental Figure S4 C–C′′′, respectively). All intensities were normalized to the intensity of the first frame (−5 s). Axotomy occurred at 0 s, indicated as the black arrowhead. Scale bar, 5 μm.

FIGURE 6: Reorganization of microtubules after axotomy. (A) Quantification of axonal regeneration percentage in control, EGTA-treated, vinblastine-treated, elav-KHC-RNAi, and ciliobrevin D-treated neurons. The y-axis represents the regeneration ratio (≥5-μm neurite regrowth overnight is counted as a regeneration event) of all the samples examined. (B, C) Microtubules of mixed polarity formed after axotomy. (B) A phase-contrast image of the neuron, with white arrowhead showing the axotomy site. (C) Kymographs of EB1-GFP comets in the segment of the axon next to the axotomy site marked with white dashed line in B: before (top), 1 min after (middle), and 25 min after (bottom) axotomy. (D–D‴) Inhibition of microtubule dynamics by 10 nM vinblastine 4 h after axotomy does not prevent Jupiter-GFP–labeled microtubule movement and looping. Looping microtubules are indicated by orange arrowheads, and sprouting microtubules are indicated by blue arrowheads. The time of adding 10 nM vinblastine is 0 min (120 min after imaging). Scale bar, 5 μm.
found that enhanced JNK activity dramatically increased neuron survival and initial axon regeneration rate (Figure 7C), consistent with published reports (Hammarlund et al., 2009; Xiong et al., 2010; Nix et al., 2011). In an opposite experiment, we blocked the JNK pathway with a JNK-specific inhibitor SP600125 (Bennett et al., 2001) and observed inhibition of both regeneration and microtubule motility in control neurons (Figure 7C and Supplemental Figure S7, A and A′) and CA JNKK neurons (Figure 7C).

We noticed that CA JNKK mature neurons (4 d in culture) usually contained more looping and bending microtubules in the cell body and fewer bundled microtubules in the axons, which implies an enhanced level of microtubule sliding. We then examined whether JNK signaling promotes microtubule sliding in mature neurons. We observed a significantly higher level of sliding in CA JNKK neurons (Figure 7D). Microtubules in CA JNKK neurons clearly move outside of the photoconversion zone and slide into neurites (Figure 7, F–F′′′′, orange arrowheads), whereas in control

**FIGURE 7:** JNK regulates neuronal regeneration by enhancing microtubule sliding. (A, B) JNK pathway activity is stimulated by axonal injury. JNK transcriptional activity is visualized by an enhancer trap of JNK target gene, puckered (puc)-lacZ, in a control intact neuron (A) and an axotomized neuron (B). (A′, B′) Boxed area in A and B, respectively. Fluorescence images in A′ and B′ were acquired and adjusted identically. White arrowhead, axotomy position. (C) Quantification of axonal regeneration percentage in constitutively active JNK mutant (CA JNKK), JNK inhibitor (SP600125)-treated, and actinomycin D–treated neurons. The y-axis represents the regeneration ratio (≥5-μm neurite regrowth overnight is counted as a regeneration event) of all the samples examined. (D) Quantification of fluorescence signal outside of the photoconversion zone after 21 min in control and CA JNKK neurons, both 4 d in culture. Unpaired t test: p < 0.005. (E–E′′′′) A 4-d control neuron has no detectable microtubule sliding. (F–F′′′′) A 4-d CA JNKK neuron slides microtubules from the cell body to neurites (indicated by orange arrowheads). Purple circle, photoconverted area; scale bar, 5 μm.
neurons, microtubules show very little movement (Figure 7, E–E‴′, and Supplemental Video S9). Therefore we conclude that JNK pathway activation stimulates microtubule sliding in mature neurons, which in turn promotes initial axonal regeneration.

JNK signaling can affect microtubule sliding and initial regeneration through two possible mechanisms: 1) JNK or another component of JNK kinase pathway could directly phosphorylate and regulate the activity of an essential regulator of microtubule sliding; 2) JNK can activate Jun/Fos-dependent transcription of a gene encoding an essential player regulating microtubule sliding. To distinguish these two mechanisms, we used a transcriptional inhibitor, actinomycin D, to stop transcriptional activity in neurons. We tested actinomycin D efficiency in cultured neurons using a RU486-inducible system (Osterwalder et al., 2001; Nicholson et al., 2008; Supplemental Figure S7B) and found that 10 μg/ml actinomycin D is sufficient to inhibit transcriptional activity in mature neurons (Supplemental Figure S7C; see details in Materials and Methods). The inhibition of transcription by actinomycin D prevents initial regeneration both in control and CA JNKK neurons (Figure 7C). The effect of actinomycin D on initial regeneration is not caused by cell lethality, because microtubules stay intact, and mitochondria are still moving in neurons after an overnight actinomycin D treatment (Supplemental Figure S7, D and D‴′, and unpublished data); this is consistent with the literature demonstrating that transcription is dispensable for survival in differentiated cultured neurons (Donady et al., 1975). Furthermore, the microtubule network before and after overnight imaging appeared static (compare Supplemental Figure S7, D‴′ to D), implying that no microtubule sliding occurred during the imaging period. Furthermore, we examined microtubule sliding rates in cultured Drosophila S2 cells (Jolly et al., 2010; Barlan et al., 2013) immediately after actinomycin D treatment and found that there was no significant difference from control (Supplemental Figure S7E). Therefore actinomycin D does not directly inhibit microtubule sliding. Together these data indicate that JNK signaling enhances microtubule sliding and axonal regeneration through transcriptional activation.

**DISCUSSION**

Numerous studies of axotomy have demonstrated the importance of calcium influx, microtubule dynamics, and JNK signaling in axonal regeneration (Liu et al., 2011). However, it is not clear how these factors integrate to drive axon regrowth. In this study, we showed that initial axon regeneration, like initial outgrowth of axons (Lu et al., 2013b), is driven by microtubule–microtubule sliding. We demonstrated that after injury, calcium influx induces localized transient microtubule disassembly at the injury site and conversion of microtubule arrays of uniform polarity to mixed-polarity arrays. Creation of these arrays allows kinesin-1 to drive microtubule sliding and push the membrane of regenerating axons, powering their initial outgrowth (see the model in Figure 8).

In this study, we observed that axotomy induces a fast calcium influx in neurons that triggers microtubule depolymerization near the injury site, which is essential for axonal regeneration. Our observation of calcium influx upon axotomy is consistent with previous studies in several other organisms, including Aplysia neurons (Ziv and Spira, 1995, 1997; Kamber et al., 2009), C. elegans sensory neurons (Ghosh-Roy et al., 2010), Drosophila sensory neurons (Ghannad-Rezaie et al., 2012), rat cortical neurons (Mandolesi et al., 2004), and mouse peripheral neurons (Cho and Cavalli, 2012; Cho et al., 2013). We further demonstrated that fast microtubule depolymerization near the injury site is induced by calcium influx, since chelation of Ca\(^{2+}\) by EGTA inhibits calcium influx and microtubule depolymerization, and results in failure of regeneration. We propose that this calcium-induced microtubule disassembly is the first step in creating microtubule arrays of mixed polarity in the distal axons, which could be the initial injury signal for neuronal regeneration.

Our description of microtubule dynamics in regenerating Drosophila axons is in agreement with previous reports that have
demonstrated dramatic reorganization of microtubules after axonal damage in several other organisms. In *Aplysia* neurons, axotomy causes reorganization of microtubules (Kamber et al., 2009), and formation of microtubule-filled filopodia-like protrusions (Goldberg and Burmeister, 1992). In *C. elegans* posterior lateral microtubule (PLM) sensory neurons, microtubule growth after axonal injury is stimulated by down-regulation of depolymerizing Kinesin-13 family member KLP-7, and persistent microtubule growth is supported by elevating A2 posttranslational modification of α-tubulin (Ghosh-Roy et al., 2012). In *Drosophila* sensory neurons, global up-regulation of microtubule dynamics and mixed microtubule orientation in dendraites have been documented after axotomy (Stone et al., 2010; Chen et al., 2012; Song et al., 2012). In mouse peripheral neurons, tubulin near the injury site has to be deacetylated by the Histone Deacetylase 5 (HDAC5) to initiate axonal regeneration (Cho and Cavalli, 2012). In rat sensory DRG (dorsal root ganglion) neurons, the kinesin-2 family member KIF3C has been implicated as a microtubule-stabilizing factor that is essential for axonal regeneration (Gumy et al., 2013).

Here we showed that formation of mixed microtubule orientation through fast microtubule depolymerization and subsequent re-polymerization are required for initial axonal regeneration. Stabilization of microtubules by treatment with a low concentration of Vinblastine prevents their depolymerization and subsequent microtubule polarity change, and suppresses regeneration, while Vinblastoine added after initial polarity orientation change does not affect regeneration. In our previous studies (Jolly et al., 2010; Lu et al., 2013b), we proposed that KHC drives microtubule-microtubule sliding by using its C-terminal tail to bind one microtubule (Hackney and Stock, 2000; Seeger and Rice, 2010), while using its processive motor domain to move along another microtubule. Since there are no signs of positive cooperatively binding of the KHC tails to microtubules in either S2 cells (our unpublished data) or in mammalian cells (Navone et al., 1992), we speculate that KHC binds to two microtubules through its motor and tail domains in a random orientation. Furthermore, a recent study from our lab has demonstrated that mitotic kinesin-6, Pav/KLP, known to cross-link antiparallel microtubules at mitotic spindles (Adams et al., 1998), downregulates microtubule sliding in both S2 cells and Drosophila neurons in vitro and in vivo (del Castillo et al., 2015). Therefore we propose that an antiparallel arrangement of microtubules is required for effective KHC sliding (Figure 8). Formation of mixed polarity microtubules in axotomized axons allows KHC to slide these microtubules apart, and this process provides the mechanical force necessary to generate membrane protrusion for initial axonal regrowth. Therefore our data not only are consistent with previous reports on microtubule destabilization, enhanced dynamics, and reorganization of polarity after axotomy, but also provide a unifying mechanism explaining the requirement for microtubule reorganization. Furthermore, the C-terminal domain of KHC involved in microtubule-microtubule sliding is evolutionary conserved across the animal kingdom (Hackney and Stock, 2000; Seeger and Rice, 2010), and general changes in microtubule dynamics are characteristic of axonal regeneration in many organisms. We propose that the microtubule-sliding mechanism uncovered in this work for initial regeneration in *Drosophila* neurons also functions in the neuronal injury response in higher organisms.

Although microtubule reorganization, induced by changes in microtubule polymerization and polarity, is required for regeneration, the reorganization alone is not sufficient to activate Kinesin-1-driven microtubule sliding and injury-induced axonal regrowth. In this study, we revealed that JNK signaling promotes microtubule sliding to provide mechanical force for initial regeneration. The JNK pathway and its upstream mitogen-activated-protein kinase (MAPK) cascade have been implicated as an essential regeneration-regulating mechanism in multiple neuronal systems, including Caenorhabditis elegans motor neurons (Hammarlund et al., 2009; Nix et al., 2011; Ghosh-Roy et al., 2012), Drosophila motor (Xiong et al., 2010; Xiong and Collins, 2012) and sensory (Stone et al., 2010; Chen et al., 2012) neurons, and adult mouse dorsal root ganglion (DRG) neurons (Itoh et al., 2009; Barnat et al., 2010). Our results indicate that JNK involvement in initial axonal regeneration is dependent on its transcriptional activation. We documented that JNK signaling increases microtubule sliding in mature neurons, and this elevated level of sliding microtubule provides mechanical force for initiation of regenerating protrusions. However, it is likely that additional factors downstream of JNK transcription activation, together with microtubule sliding, contribute to long-range axonal regeneration (Figure 8). It has been shown that JNK signaling promotes stem cell maintenance in a transcription-dependent manner in *Drosophila* (Lu et al., 2012), and therefore it is tempting to propose that JNK signaling could reprogram the mature neurons back to a less-differentiated status through transcriptional activation to reinitiate neurite out-growth. Further studies on JNK transcriptional target genes involved in microtubule sliding and examination of JNK-promoting microtubule sliding in neurons of higher organisms will provide more insight into the mechanism of initial axonal regeneration, as well as aid in the development of effective therapeutic treatments for spinal cord injury and brain damage.

**MATERIALS AND METHODS**

**Drosophila genetics**

Fly stocks and crosses were kept on regular cornmeal food supplemented with dry active yeast at room temperature (~22°C) and switched to apple juice agar food supplemented with dry yeast 2–3 d before embryo collection. For larval motor neuron regeneration experiments, D42-Gal4 (Pilling et al., 2006) was used to specifically drive UAS expression in motor neurons. For embryonic neuron regeneration experiments, elav-Gal4 (Campos et al., 1987; Robinow and White, 1988) was used to drive UAS expression in the nervous system. To knock down KHC in larval brain neurons, transheterozygote Khc027/Khc027 flies (Brendza et al., 1999, 2000; Serbus et al., 2005) were used. To knock down DHC in larval brain neurons, a TRiP RNAi line (Bloomington stock #36583, encoding a short RNA hairpin against the DHC64C CDS 10044-10064) was driven by elav-Gal4. To knock down KHC in mature embryonic neurons, a TRiP RNAi line (Bloomington stock #35770, encoding a short RNA hairpin against the KHC 3’ UTR 231–251) was driven by elav-Gal4.

**Larval motor neuron culture**

The protocol is modified based on the protocol from the Sprecher lab (Egger et al., 2013). *Drosophila* third-instar larvae were picked and washed with distilled H2O, 70% EtOH, and sterile 1x modified dissection saline (9.9 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid, pH 7.5, 137 mM NaCl, 5.4 mM KCl, 0.17 mM NaH2PO4, 0.22 mM KH2PO4, 3.3 mM glucose, 43.8 mM sucrose). Optic lobes and ventral ganglion were dissected and cleaned in sterile 1X modified dissection saline; tissues were then dissociated in Liberase solution (Roche, Basel, Switzerland; Liberase TM Reagent, 0.02 mg/ml 13 Wünsch units/ml in sterile 1X modified dissection saline), final concentration in 0.25 mg/ml 1.3 Wünsch units/ml for 1 h. The neurons were spun down at 300 × g for 5 min and then washed twice in supplemented...
Schneider's medium (20% fetal bovine serum, 5 μg/ml insulin, 100 μg/ml penicillin, 100 μg/ml streptomycin, and 10 μg/ml tetracycline); the neurons were plated onto concanavalin A–coated coverslips and allowed attached for 30 min before addition of full volume of supplemented Schneider’s medium for imaging 2–4 h after plating or incubated for 2–3 d before imaging.

Embryonic neuronal axotomy procedure

The detailed embryonic neuronal preparation protocol is available online (www.jove.com/video/50838/organelle-transplant-cultural-drosophila-cells-s2-cell-line-primary; Lu et al., 2013a). Cultured embryonic neurons were kept in 25°C in an insect cell incubator for 4–5 d in supplemented Schneider’s medium with 5 μM cytochalasin D; cytochalasin D was then washed out and 50 μg/ml gentamicin added for the axotomy experiment. A fine glass needle was prepared from a borosilicate glass capillary (outer diameter, 1.00 mm; inner diameter, 0.78 mm) using a Flaming/Brown Micropipette Puller (model P-97, Sutter Instrument, Novato, CA) and manually controlled by a Nikon Narishige (Tokyo, Japan) micromanipulator. The needle tip usually scratches against the axon perpendicularly once or twice to complete the axotomy. As indicated, the following drugs were added to cultured neurons before the axotomy procedure: EGTA (10 mM), vinblastine (10 nM), ciliobrevin D (20 μM), SP600125 (20 μM), and actinomycin D (10 μg/ml).

Microscope image acquisition

A Nikon (Tokyo, Japan) Eclipse U2000 inverted microscope equipped with S-Flor 40x/1.30 numerical aperture (NA) and Plan-Apo 100x/1.4 NA objectives was used to perform neuronal axotomy and image regeneration. Images were acquired by a digital complementary metal–oxide–semiconductor camera, ORCA-Flash4.0 V2 C11440-22CU (Hamamatsu Photonics, Hamamatsu, Japan) controlled by MetaMorph software, version 7.7.7.0 (Molecular Devices, Sunnyvale, CA). Phase-contrast images were obtained using a 100-W halogen lamp or CoolLED PrecisExcite (Hampshire, UK), and fluorescence excitation was achieved using an Aura light engine (Lumencor, Beaverton, OR) or X-Cite XLED (Lumen Dynamics, Mississauga, Canada). tdEOS- etub neurons were photoconverted by ultraviolet light from a mercury light source through a pinhole in the light path for 3 s. TIRF images after photoconversion were acquired with a Plan-Apo TIRF 100x/1.45 NA lens and a Sapphire 561-nm laser (Coherent, Santa Clara, CA). A Nikon Eclipse U2000 inverted stand with a Yokogawa CSU10 spinning disk confocal head and a 100x/1.45 NA lens were used to image nonaxotomy samples. Images were acquired using an Evolve 512 electron-multiplying charge-coupled device camera (Photometrics, Tucson, AZ) controlled by Nikon NIS-Elements software (AR 4.0.07 64bt). Phase-contrast images of tdEOS- etub were acquired using illumination from a Heliophor 89 North light (Burlington, VT) in the epifluorescence pathway by a 405-nm filter and confined by an adjustable pinhole in the field diaphragm position (projected as an 8–12-μm-diameter circle in the sample plane) for 6 s. All neurons were imaged in 20% fetal bovine serum–supplemented Schneider’s medium (Sigma-Aldrich, St. Louis, MO), except for the ciliobrevin D treatment experiments, for which imaging was in nonsupplemented Schneider’s medium. S2 cells were imaged in Gibco Insect Xpress medium (Life Technologies, Carlsbad, CA). All images were acquired in 16-bit format at room temperature (~24°C), analyzed by Fiji/ImageJ, and plotted in GraphPad Prism 5. Heat-map images of the calcium reporter (GCaMP3) were created using the ImageJ Heatmap Histogram plug-in (www.samuelpean.com/heatmap-histogram/). tdEOS- etub848B images were bleach corrected using the ImageJ Bleach Correction plug-in (fiji.sc/Bleach_Correction). All figures were assembled in Adobe Illustrator CS4, and all videos were assembled in AVS Video Editor 5.1.

Immunocytochemistry for puc669-lacZ

Embryonic neurons from the embryos of puc669-lacZ/TM6B were plated on concanavalin A–coated, 25-mm photoetched coverslips (#72265-25; Electron Microscopy Sciences, Hatfield, PA) for >48 h. Neurons were first injured by a glass needle (see Embryonic neuronal axotomy procedure for more details) with the position information recorded according to the grids. After axotomy, neurons were cultured overnight (>16 h) to allow the expression of the reporter. Then neurons were washed three times in 1× phosphate-buffered saline (PBS) before fixation in 4% formaldehyde for 10 min at room temperature and then washed three times in 1× PBS and washed and blocked in wash solution (0.1% Triton X-100, 1% bovine serum albumin in Tris-buffered saline) for 10 min three times. Then they were incubated with the primary antibody, a monoclonal anti–β-galactosidase antibody (1:250; Cat# Z3781; Promega, Madison, WI), for 45 min at room temperature in a moisture chamber, washed and blocked in wash solution 10 min three times, incubated with secondary antibody, goat anti-mouse tetramethylrhodamine isothiocyanate (1:200; Jackson Immuno-Research, West Grove, PA) for 45 min to 1 h in the dark at room temperature, and washed and blocked in wash solution 10 min three times. Cells were covered with 1× PBS and imaged directly according to the grid position information.

Quantification of fluorescence intensity of microtubules

Time-lapse movies of elav>tdEOS- etub neurons (X-Cite 460 channel) were aligned using the StackReg plug-in (fiji.sc/StackReg). Same-size regions of interest (ROIs) were assigned to different positions using ROI Manager, and the fluorescence intensity of designated ROIs over time (multimeasure in ROI Manager) was measured. Background noise was subtracted from the raw data, and the intensities of all frames were normalized according to the intensity of the first frame. The data were analyzed and plotted in GraphPad Prism 5.

Test of actinomycin D efficiency

Four identical samples of primary embryonic neurons were prepared from embryos expressing elav>GeneSwitch>mCD8-GFP. Sample 1 was treated with only dimethyl sulfoxide. Samples 2–4 were bathed in 50 μg/ml RU486 for 48 h before washing three times by supplemented Schneider’s medium. Samples 3 and 4 were then treated overnight with 10 μg/ml actinomycin D, respectively. After 18 h of incubation, samples 1–4 were imaged to determine the percentage of GFP-positive neurons.

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