Facilitation of axon regeneration by enhancing mitochondrial transport and rescuing energy deficits

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Although neuronal regeneration is a highly energy-demanding process, axonal mitochondrial transport progressively declines with maturation. Mature neurons typically fail to regenerate after injury, thus raising a fundamental question as to whether mitochondrial transport is necessary to meet enhanced metabolic requirements during regeneration. Here, we reveal that reduced mitochondrial motility and energy deficits in injured axons are intrinsic mechanisms controlling regrowth in mature neurons. Axotomy induces acute mitochondrial depolarization and ATP depletion in injured axons. Thus, mature neuron-associated increases in mitochondria-anchoring protein syntaphilin (SNPH) and decreases in mitochondrial transport cause local energy deficits. Strikingly, enhancing mitochondrial transport via genetic manipulation facilitates regenerative capacity by replenishing healthy mitochondria in injured axons, thereby rescuing energy deficits. An in vivo sciatic nerve crush study further shows that enhanced mitochondrial transport in snph knockout mice accelerates axon regeneration. Understanding deficits in mitochondrial trafficking and energy supply in injured axons of mature neurons benefits development of new strategies to stimulate axon regeneration.

Introduction

Mitochondria are cellular power plants that supply ATP essential for neuron growth, survival, and regeneration (Nicholls and Budd, 2000). Because of their extremely varied morphological features, neurons face special challenges to maintain energy homeostasis in distal regions of the axons. Mitochondrial distribution to far distal axons depends on microtubule (MT)-based motors via mechanisms that require ATP hydrolysis (Ruthel and Hollenbeck, 2003; MacAskill and Kittler, 2010; Saxton and Hollenbeck, 2012; Sheng and Cai, 2012). Mitochondria move bidirectionally along axons and frequently change direction. Motile mitochondria can become stationary, and stationary ones can be redistributed in response to changes in metabolic status and growth conditions (Sheng, 2014). In mature axons of the central nervous system (CNS), the majority of mitochondria remain stationary, whereas ~20–30% are motile (Sun et al., 2013). In addition, distal mitochondria need to be removed when their integrity is impaired under certain stress conditions (Miller and Sheetz, 2004; Chang and Reynolds, 2006; Cai et al., 2012).

Although young neurons possess the capacity for robust axon growth during early development, mature CNS axons typically fail to regrow after injury, leading to permanent neurological impairments. Numerous studies in the past have focused on genetic programs, signaling mechanisms, and extracellular inhibitory factors for axon regeneration (see reviews by Schwab and Bartholdi [1996], Case and Tessier-Lavigne [2005], Filbin [2006], Harel and Strittmatter [2006], Yiu and He [2006], Fitch and Silver [2008], Liu et al. [2011], and Cho and Cavalli [2014]). These studies suggest that mature CNS neurons have lost their growth capacity as the result of an intrinsic decline of permissive conditions for regeneration. Thus, it is critical to reveal these intrinsic pathways that account for mature neuron-associated decline of axonal regrowth capacity.

To survive an injury, neurons need to quickly reform an active growth cone where damaged membranes are resealed, cytoskeletal structures are rearranged, and regrowth programs are activated, including synthesis of raw materials, transport, and assembly of axonal components (Bradke et al., 2012). All of these events require high levels of energy consumption. Thus, proper mitochondrial transport into injured axons ensures that metabolically active areas are adequately supplied with ATP. However, two fundamental questions remain to be addressed. (1) Do mature neurons maintain an effective capacity to recruit mitochondria to injured axons? And (2) if this function declines in mature neurons, does enhancing mitochondrial transport enable mature neurons to regain regenerative capacity.
by facilitating the removal of damaged mitochondria from and replenishing healthy ones to injured axons?

Our previous study demonstrated that syntaphilin (SNPH) acts as a “static anchor” specific for axonal mitochondria (Kang et al., 2008; Chen and Sheng, 2013). Deleting murine snph results in a substantially increased percentage (~70%) of motile axonal mitochondria. Conversely, overexpressing SNPH abolishes axonal mitochondrial transport. Most interestingly, SNPH is strictly developmentally regulated in the brain: its expression is hardly detectable in embryonic stages, very low before post-natal day 7 (P7), and peaks at adult stages (Das et al., 2003). The unique pattern of SNPH expression in the brain and its specific role in anchoring axonal mitochondria allow us to propose an attractive hypothesis: mature neuron-associated decline of mitochondrial transport is an intrinsic mechanism controlling axon regrowth capacity. Thus, snph knockout (KO) mice provide an ideal model to investigate how mitochondrial trafficking and anchoring influences axonal regenerative capacity.

In the current study, we reveal that enhancing axonal mitochondrial transport in snph KO mice facilitates axon regenerative capacity. We further show that axon injury is an acute stress signal that depolarizes mitochondria in the vicinity of the injured site. Enhancing transport helps remove those dysfunctional mitochondria from and replenish healthy ones to the injured axons, thus rescuing energy deficits and facilitating axonal regrowth. Therefore, our study suggests a new cellular target for stimulating axon regeneration and perhaps functional recovery after nerve injury and disease.

**Results**

**Enhanced axon regrowth in snph KO cortical neurons**

Our previous study identified SNPH as an axonal mitochondrial anchoring protein; deleting snph results in a majority (~70%) of motile axonal mitochondria in cultured hippocampal and cortical neurons (Kang et al., 2008; Chen and Sheng, 2013). To examine the in vivo effect of SNPH on axonal mitochondrial transport, we crossed snph KO mice with Thy1-Mito-CFP transgenic mice, where mitochondria are labeled with CFP in a subset of neurons (Misgeld et al., 2007). Relative mitochondrial motility was assessed from acute nerve explants after rapid dissection in adult wild-type (WT) and snph KO littermates (2 mo old). The crossed snph−/+ /Mito-CFP mice display robustly enhanced mitochondrial motility along axonal bundles of the sciatic nerve (71.58 ± 2.72%, mean ± standard error, P < 0.0001, Mann–Whitney U test) relative to that in control Mito-CFP mice (30.95 ± 2.17%, n = 23 images for each genotype from five paired littermates; Videos 1 and 2). This ex vivo imaging further verifies that SNPH acts as an anchoring protein that controls axonal mitochondrial transport. Thus, snph KO mice serve as an ideal genetic model for our investigations as to how enhanced transport of axonal mitochondria in mature neurons influences axonal regrowth and regenerative capacity.

We first asked whether enhanced mitochondrial transport in snph KO cortical neurons facilitates axon regrowth after axotomy. To address this issue, we applied microfluidic culture devices as we previously described (Zhou et al., 2012), in which neuronal cell bodies and dendrites are restricted to the soma chamber while axons grow into the axon terminal chamber through long (450 μm in length) microgrooves. Neurons at 4 d in vitro (DIV4) were stained with βIII-tubulin, which labels axons (Fig. 1 A). Deleting snph in cortical neurons did not change axonal growth during the early stages of development (DIV5; P = 0.38, n = 62) when compared with WT neurons (n = 60; Fig. S1, A and B). However, when axons in the terminal chamber were removed by vacuum aspiration (axotomy; Fig. 1 B), we found that snph KO neurons display enhanced regrowth capacity 3 d after axotomy (Fig. 1, C–E). Axon regrowth begins as early as 7 h after injury and enters a rapid growth phase by 28 h (Fig. 1, F and G). Regrown axonal areas are significantly increased at 28 h in snph KO neurons relative to WT control (P < 0.001). We costained terminal axons with phalloidin to label active growth cones 14 h after axotomy. Deleting snph significantly facilitates formation of new growth cones (69.11 ± 2.90%, P = 0.0002) compared with WT neurons (44.46 ± 1.92%; Fig. 1, H and I). Live imaging shows that regenerating axons are mainly grown from the injured tips (Fig. S1 C). Thus, enhancing mitochondrial transport by deleting the snph gene facilitates axon regrowth after injury.

**Mature neuron-associated decline of mitochondrial transport contributes to failed regrowth**

We asked whether SNPH expression contributes to a reduced regrowth capacity in mature CNS neurons. WT cortical neurons in microfluidic devices were axotomized at DIV4 or DIV12 and stained with βIII-tubulin 3 or 6 d after axotomy. Although neurons at DIV7 maintain some regrowth capacity, mature neurons at DIV18 display little axon regrowth (Fig. 2, A–C). A possible mechanism underlying such a striking difference in regrowth capacity between young and mature neurons may be caused by development-dependent cellular pathways. To address this issue, we examined the relative expression of SNPH and other mitochondrial proteins in various developmental stages of cultured neurons between DIV3 and DIV22, mouse brains at embryonic day (E) 18 and adult. To our surprise, mature neuron-associated decline of regrowth capacity correlates with progressively increased SNPH expression: its expression is hardly detectable in E18 brains and in cultured neurons before DIV3, readily detected after DIV9, and peaks at DIV22 in culture (Fig. 2, D and E), an expression pattern consistent with our previous findings from rat brains (Das et al., 2003). There were no other proteins detected showing such changes, including kinesin-1 (KIF5), kinesin adaptors Trak2 and Miro1/2, dynein intermediate chain (DIC), and outer mitochondrial membrane proteins TOM20 and VDAC. Furthermore, there is no significant change (P = 0.25) in Miro1/2 expression in WT and snph KO cortical neurons at DIV14 (Fig. S1 D). In addition, we expressed HA-SNPH or HA control in cortical neurons and detected no observable change in Miro1/2 expression at DIV14 (P = 0.16; Fig. S1 E).

The unique expression pattern suggests that SNPH controls mitochondrial transport in mature neurons. Consistent with this hypothesis, axonal mitochondria display progressively reduced motility from DIV7 to DIV18 (Fig. 2, F and G). The relative axonal mitochondrial motility at DIV7 is 47.60 ± 4.60%, two times higher than that at DIV18 (22.88 ± 3.09%, P < 0.001). To exclude a general transport decline caused by the long-term neuronal culture, we examined the axonal transport of late endosomes as an internal control and found no decline during the same maturation stages (Fig. 2, F and H). These results suggest that elevated SNPH expression in mature neurons is one of the intrinsic mechanisms diminishing axonal regenerative capacity.
Mature neurons regain regrowth capacity by enhancing mitochondrial transport

Mature neurons at DIV18 lose their regrowth capacity after axotomy (Fig. 2, A–C) and display a substantial decline in axonal mitochondrial transport (Fig. 2, F–H), raising a question as to whether mature neurons can regain their regrowth capacity by enhancing mitochondrial transport. To address this issue, we manipulated axonal mitochondrial motility in cortical neurons by coexpressing DsRed-Mito with SNPH, SNPH-dMTB, an SNPH loss-of-function mutant deleting its anchoring domain (Kang et al., 2008), or Miro1 via lentivirus infection. Cultured neurons display strikingly different motility patterns of mitochondria along the microgrooves after transgene expression (Fig. 3 A). Expressing SNPH abolishes axonal mitochondrial motility, whereas expressing Miro1 enhances their bidirectional transport along the microgrooves relative to control neurons (Fig. 3 B and Videos 3–5). To assess axon regrowth after axotomy, these cortical neurons were coinfected with Lenti-GFP to visualize axons in the terminal chambers and grown for 12 d before axotomy. Axonal regrowth was evaluated 6 d after injury (DIV18). Although neurons coexpressing SNPH showed very limited axon regrowth, enhancing mitochondrial transport by expressing Miro1 robustly increases axon regrowth capacity when compared with neurons expressing SNPH-dMTB (Fig. 3, C and...
D). SNPH-mediated suppression of axon regrowth is partially recovered by supplying 200 µM ATP through electroporation (Fig. 3 E). To determine whether mitochondrial ATP production is critical for axon regrowth, we briefly treated the axonal chamber with 2 µM oligomycin, a mitochondrial complex V inhibitor, after axotomy. Such treatment abolishes recovery of the regrowth capacity in snph KO neurons (Fig. 3 F). Altogether, these results suggest that supplying ATP to injured axons by enhancing mitochondrial transport contributes to axon regrowth.

To determine whether increased regrowth in snph KO neurons is secondary to improved neuronal survival, we examined MAP2-positive survival neurons and TUNEL-positive death cells in the somatic chamber (Fig. S2 A) and characterized the correlation between cell density and axonal regrowth after injury. Our study shows no significant change in the normalized density of MAP2-positive neurons (Fig. S2 B) and percentage of TUNEL-positive death cells (Fig. S2 C) between WT and snph KO neurons at 1 h and 3 d after axotomy (P > 0.05). Although WT neuron density correlates with axonal regrowth after injury (R² = 0.3501, P < 0.0001), enhanced axonal regrowth in snph KO neurons is not correlated with relative cell density in the soma chamber (R² = 0.0556, P = 0.1187; Fig. S2 D). Our results are consistent with previous studies showing that promoting neuronal survival failed to increase axon regeneration (Park et al., 2008; Hu et al., 2012).

To further test our hypothesis, we examined the regrowth capacity of mature dorsal root ganglion (DRG) neurons isolated from adult WT or snph KO mice at postnatal day (P) 60. These adult DRG neurons are well matured and undergo an axotomy-like injury during cell dissociation and culture preparation; axon regrowth occurs from the cell body 1 h after plating on coverslips, thus partially recapitulating an axonal regrowth program in mature neurons after injury. Because almost all processes in DRG neurons are tau-positive axons (Perlson et al., 2009), adult DRG neurons were immunostained with the neuron-specific marker βIII-tubulin. The snph KO adult DRG neurons display increased regrowth capacity, as indicated by the total number of axonal intersections when compared with WT.
DRG neurons (P < 0.05; Fig. 3, G and H), thereby confirming that axonal regrowth in matured neurons is facilitated by enhancing mitochondrial transport.

**Expressing SNPH alters mitochondrial distribution and ATP/ADP ratio in distal axons**

A critical step for inducing axon regrowth is the transformation of the injured axonal end into a growth cone–like structure that can integrate extracellular and intracellular signals for regrowth (see reviews by Tessier-Lavigne and Goodman [1996], Yu and Bargmann [2001], and Liu et al. [2011]). We next addressed whether altered mitochondrial motility regulates their distribution in distal axons, thus controlling the size of growth cones in both the peripheral nervous system (PNS) and CNS. We first used adult DRG sensory neurons isolated from adult P60 mice. Neurons were cotransfected with DsRed-Mito with GFP and SNPH, SNPH-dMTB, or Miro1, followed by immunostaining and quantification of mitochondrial motility and axonal regrowth. Axon regeneration was evaluated 6 d after axotomy (DIV18). Note that abolishing mitochondrial transport by expressing SNPH shows failed axon regrowth, whereas enhancing mitochondrial transport by expressing Miro1 robustly increases axon regrowth capacity.

**Figure 3.** Mature neurons regain regrowth capacity by enhancing mitochondrial transport. (A and B) Kymographs (A) and quantitative analysis (B) showing mitochondrial motility along microgrooves in microfluidic chambers. Time-lapse imaging was recorded in cortical neurons at DIV12 for a total of 100 frames with 5-s intervals. In kymographs, vertical lines represent stationary organelles; oblique lines or curves to the right indicate anterograde transport toward distal terminals. Note that the relative motility in control neurons expressing HA is significantly higher than the motility in neurons overexpressing HA-SNPH, but lower than the motility in neurons expressing HA-Miro1. [C and D] Representative microfluidic images (C) and quantitative analysis (D) showing regrowth capacity in mature cortical neurons. Neurons infected with lentivirus encoding SNPH, SNPH-dMTB, or Miro1 were grown on microfluidic chambers for 12 d before axotomy. Axon regeneration was evaluated 6 d after axotomy (DIV18). Note that abolishing mitochondrial transport by expressing SNPH shows failed axon regrowth, whereas enhancing mitochondrial transport by expressing Miro1 robustly increases axon regrowth capacity. (E) Partial recovery of regrowth capacity in the SNPH-expressing neurons by ATP application. The electroporated neurons were immediately plated on a microfluidic chamber with medium containing 200 µM ATP. [F] Recovery of regrowth capacity 1.4 h after axotomy in snph KO neurons is largely abolished by blocking mitochondrial ATP generation with 2 µM oligomycin (Oligo). The axonal chambers were briefly treated with 2 µM oligomycin for 4 h after axotomy. (G and H) Representative images (G) and quantitative analysis (H) showing enhanced axonal regrowth in snph KO adult DRG neurons. DRG neurons isolated from adult (P60) WT or snph KO mice were immunostained with βIII-tubulin at DIV1. Axon regrowth was quantified by Sholl analysis. The snph-deficient adult DRG neurons display increased axon branching as indicated by the total number of axonal intersections. Mitochondrial motility data were analyzed from the total number of microgrooves (B); axonal regrowth data were analyzed in terminal chambers where new axons grow from a total number of microgrooves (E and F), total number of terminal chambers (D), or total number of DRG neurons (H) indicated within bars or in parentheses and expressed as mean ± SE and by one-way ANOVA test (B and D), Student’s t test (F and H), or Mann–Whitney U test (E). Bars: (A) 20 µm; (C and G) 100 µm.
with the axonal marker Tau. Expressing SNPH, but not its loss-of-function mutant dMTB, blocks delivery of mitochondria into axonal terminals (Fig. 4, A and B). Conversely, expressing a loss-of-function mutant SNPH-dMTB, blocks the delivery of mitochondria into growth tips and reduces average size of growth cones, whereas expressing Miro1 robustly increases terminal mitochondrial density and size of growth cones. C and D) Images (C) and quantitative analysis (D) of distal mitochondrial distribution and growth cone size in cortical neurons coexpressing DsRed-Mito with GFP or together with SNPH, SNPH-dMTB, or Miro1. Neurons at DIV5 were immunostained with Tau. Note that both mitochondrial density in terminals and the average size of growth cones are decreased in neurons expressing SNPH (P < 0.01 and P < 0.001, respectively), but increased in neurons expressing Miro1 (P < 0.05) relative to control neurons. Arrows indicate the most distal mitochondrion in the SNPH-expressing axons (C). Data were analyzed from the total number of axons indicated within bars from more than three experiments and expressed as mean ± SE and by one-way ANOVA test.

We further confirmed this correlation in cortical neurons by coexpressing DsRed-Mito with GFP or together with SNPH, SNPH-dMTB, or Miro1, followed by immunostaining with Tau. Both mitochondrial density in axonal terminals and the mean size of growth cones were decreased in neurons overexpressing SNPH (P < 0.01 and P < 0.001, respectively), but increased in neurons overexpressing Miro1 (P < 0.05) relative to control neurons (Fig. 4, C and D). Because SNPH is hardly detectable before DIV7, it is predictable that expressing loss-of-function mutant SNPH-dMTB in young cortical neurons (DIV5) displays no dominant-negative effect on mitochondrial density nor the mean size of growth cones when compared with control neurons (Fig. 4 D). Thus, our results provide a clue as to why an injured axonal tip in mature neurons often fails to transform into an active growth cone (Li and Raisman, 1995; Hill et al., 2001), as mitochondrial recruitment is suppressed by elevated SNPH expression.

Reduced mitochondrial density within axonal terminals in the SNPH-expressing neurons prompted us to examine how mitochondrial motility impacts ATP homeostasis throughout an axon. The cellular ATP/ADP ratio in live axons was measured by applying an engineered fluorescent ATP sensor, PercevalHR, in live neurons (Sun et al., 2013; Tantama et al., 2013). The fluorescence intensity ratio (F_{488nm}/F_{405nm}) reflects the relative ATP/ADP ratio, thus allowing spatial detection of intracellular ATP homeostasis in various axonal segments. Relative mitochondrial density and ATP/ADP ratios along an axon were analyzed based on five equally divided segments of each individual axon from the most proximal to the most distal region (Fig. S3, A and B). Overexpressing SNPH in cortical neurons redistributes axonal mitochondria to the proximal region (P < 0.001) and also reduces mitochondrial density and the ATP/ADP ratio in the most distal axon segment (P < 0.001 and P < 0.05, respectively). Because intracellular pH values impact the Perceval fluorescent intensity (Berg et al., 2009), we monitored pH in axons using the pH dye SNARF-5F (Fig. S3, C and D). SNARF-5F signal was calibrated using various buffered solutions containing high K+/nigericin with varying pH values (Fig. S3, E and F). The pH values in an entire axon of both WT and SNPH overexpression neurons undergo very minor changes (7.10–7.15). Thus, reduction in the ATP/ADP ratios in distal axons is physiologically relevant to the reduced axonal mitochondrial density. These results suggest that proper mitochondrial positioning in distal axons is required to maintain local ATP homeostasis. Therefore, in mature neurons, where SNPH expression is elevated and axonal mitochondrial motility is suppressed (Fig. 2), local ATP deficits occur if injury stress triggers mitochondria dysfunction. This notion is supported by previous in vivo studies showing that axonal injury induces mitochondrial depolarization and oxidative stress (O’Donnell et al., 2013; Cavallucci et al., 2014).
described (Cho et al., 2013) with some modifications. A train of near-infrared femtosecond laser pulses lead to nearly 100% physical separation of axonal processes. Time-lapse images demonstrate that although mitochondria at the axotomized site were ablated during axotomy, mitochondria in the vicinity suddenly shrank and lost their staining by tetracyanomethylene ethylester (TMRE), a fluorescent dye that stains mitochondria depending on membrane potential (Δψm, Fig. 5, A–C).

Second, we confirmed the axotomy-induced depolarization of local mitochondria in mature cortical neurons on microfluidic chambers. Neurons were infected with lentiviruses expressing pLenti-GFP or pLenti-GFP-Mito. Axon bundles in the terminal chambers were loaded with 25 nM TMRE dye at DIV12, followed by laser-based axotomy and time-lapse imaging. The axons were quickly broken up during axotomy, and mitochondria in the vicinity suddenly lost their TMRE staining (Fig. 5 D and Video 6). Similarly, in the axons expressing GFP-Mito, axotomy triggered a loss of TMRE staining near the axotomy site, while those depolarized mitochondria still maintained GFP-Mito signals (Fig. 5 E and Video 7). Alternatively, we applied a physical axotomy in the axonal chamber by using pulled glass capillaries with a tip diameter of ~1 μm. Neurons were loaded with both TMRE (Δψm-dependent dye) and MitoTracker Green FM (Δψm-independent dye) before axotomy. For physical axotomy, TMRE staining was lost near the axotomized ends, whereas MitoTracker Green FM staining remained (Fig. S4 A). In addition, we costained mitochondria in the axonal chambers with MitoTracker Green FM and MitoTracker Red FM (Δψm-insensitive dye with similar wavelength as TMRE). Laser-induced axotomy does not reduce the signal intensity of MitoTracker Red FM (Fig. S4 B), thus excluding an artificial laser effect on TMRE signals.

Third, we examined somatic mitochondrial integrity by staining with MitoTracker Orange CMTMRos, a fixable Δψm-dependent dye. Axotomy in the axonal terminal chamber does not induce any observable reduction in the mean intensity of Orange CMTMRos in the soma chamber at 1 and 5 h after injury (Fig. S4 C). In addition, we asked whether the observed local mitochondrial damage is accompanied by ATP depletion in the injured axons. To monitor intracellular ATP levels, we applied the red-shifted genetically encoded ATP probe GO-ATeam2, in which GFP and OFP were used as a FRET (Förster resonance energy transfer) pair (Nakano et al., 2011). This ATP probe is relatively stable against acidification and less phototoxic to cells and thus is suitable for long-time live imaging. Laser-based axotomy triggers a quick ATP depletion at millimolar levels in the vicinity of the injured site of adult DRG neurons (Fig. 5, F and G), indicating axotomy-induced energy deficits. These studies consistently support the notion that axotomy is an acute stress signal that depolarizes local mitochondria, thus triggering energy deficits in the vicinity of injury sites.

**Enhanced transport recovers mitochondrial integrity and rescues energy deficits**

Depolarized mitochondria not only supply less ATP, but also release toxic reactive oxygen species that trigger axon degeneration (Sheng and Cai, 2012). Therefore, quickly replacing those damaged mitochondria not only protects axons from degeneration, but also supports high-energy demanded regeneration. We hypothesize that the intrinsic reduction of mitochondrial transport in mature neurons impairs efficient delivery of healthy mitochondria to and removal of damaged mitochondria from injured axons. Mature cortical neurons can regain axon regrowth capacity by enhancing mitochondrial transport (Fig. 1, C–E; and Fig. 3 C), supporting that proper axonal transport is essential to maintain mitochondrial integrity and energy supply in response to injury-induced mitochondrial damage.

To test our hypothesis, we performed the following three types of experiments. First, we characterized axonal mitochondrial flux along the microgrooves before and after axotomy in WT and snph KO mature cortical neurons infected with pLenti-GFP-Mito. Mitochondrial flux was measured by the total number of bidirectional transport events through the microgrooves. Deleting snph significantly increases axonal mitochondrial flux before (P < 0.001) and 10 min (P < 0.05) and 1 h after axotomy (P < 0.001) when compared with WT neurons (Fig. 6, A and B; and Videos 8 and 9). Our findings also suggest that the immediate pause of axonal mitochondria upon axotomy (Videos 6 and 7) is likely a temporary response to high calcium influx into injured axons; paused mitochondria are then recruited to mobile pools 10 min after axotomy, and their total flux rate is affected by the relative SNPH expression in mature neurons.

Next, we investigated the impact of enhanced transport on the recovery of mitochondrial integrity by examining mitochondrial Δψm after injury. Cortical neurons were infected with pLenti-GFP-Mito and axons were loaded with 25 nM TMRE dye before or 1 or 5 h after axotomy. Axotomy impairs mitochondrial integrity by depolarizing Δψm in the proximal ends of the injured axons in WT neurons (Fig. 5, F and G). However, there was a significant increase (P < 0.001) in TMRE recovery in snph KO neurons (Fig. 6 D), suggesting that the mitochondrial stress phenotype can be reversed in snph KO axons by enhanced mitochondrial transport. These results support our hypothesis that enhancing mitochondrial transport helps remove damaged mitochondria and/or replenish healthy ones to the proximal ends of injured axons.

We next asked whether local ATP deficits occur after axotomy-induced mitochondrial dysfunction in mature neurons, where SNPH expression is elevated and axonal mitochondrial motility is suppressed (Fig. 2). We measured ATP maintenance in the distal growing tips 6 h after axotomy in WT and snph KO neurons infected with pLenti-PercevalHR. Enhanced mitochondrial transport in snph KO neurons significantly increases the ATP/ADP ratio in the growing tips relative to that in WT neurons (Fig. 6, E and F). We detected no change in pH values at growing tips 6 h after axotomy between WT and snph KO neurons (Fig. S5 A). To further confirm these observations, we alternatively applied the ATP probe GO-Ateam2 to monitor intracellular ATP levels by measuring the F560nm/F510nm ratio-metric integrated intensity before or 1, 3, or 5 h after axotomy. ATP-insensitive GO-Ateam3 mutant (R122K/R126K; Nakano et al., 2011) was also applied as a control of imaging and axotomy procedures. WT neurons expressing the ATP probe GO-Ateam2 display reduced F560nm/F510nm integrated intensity in the distal microgrooves after axotomy, suggesting an axotomy-induced ATP depletion. Enhancing axonal mitochondrial transport in snph KO neurons recovers energy deficits at 3 and 5 h after axotomy (Fig. 6, G and H; and Fig. S5 B). As a control, WT neurons expressing ATP-insensitive GO-Ateam3 mutant do not show any significant change after axotomy (Figs. S5, C and D). Thus, both ATP sensors Perceval and GO-Ateam2 consistently reveal a higher ATP/ADP ratio in the regrowing axon of snph KO neurons than in those of WT neurons after axotomy. We further confirmed that enhancing mitochondrial transport
Figure 5. Axotomy depolarizes mitochondria in the vicinity of injured sites. (A) Representative images showing axonal mitochondria in adult DRG neurons before and after laser-based axotomy. Adult DRG neurons isolated from P60 mice were cotransfected with DsRed-Mito and GFP. Note that mitochondria at the axotomized site were immediately lost during axotomy (white arrows). (B and C) Kymographs showing axonal mitochondria labeled by DsRed-Mito (B) or colabeled by GFP-Mito and TMRE (C) before and after axotomy. The white arrows indicate laser-scanning sites (injured site), and the black arrows in the y axis show the laser execution time. Time-lapse images were captured to show that mitochondria in the vicinity suddenly shrunk (B) and lost the $\Delta\psi_m$ (TMRE staining; C). The images were first recorded at 5-s intervals for a total of 50 frames; the consecutive post-axotomy recording was collected at 5-s intervals for a total of 50 frames. (D and E) Representative images showing axotomy-induced depolarization of axonal mitochondria. Neurons were infected with pLenti-GFP (D) or GFP-Mito (E), and axons in the terminal chambers were loaded with 25 nM TMRE dye at DIV12, followed by laser-based axotomy and time-lapse imaging. Note that in neurons expressing GFP (D), axons were quickly broken up upon axotomy (white dashed lines), and a majority of mitochondria in the vicinity suddenly lost their TMRE staining (bottom right). In neurons expressing GFP-Mito (E), axotomy triggered a sudden loss of mitochondria staining by TMRE near the axotomy site (white dashed lines), whereas those depolarized mitochondria maintained GFP-Mito signals. MC, microgroove channels. (F and G) Representative images (F) and ratiometric kymograph (G) showing axotomy-induced ATP depletion. Cultured adult DRG neurons from 2-mo-old mice were transfected with red-shifted ATP probe GO-ATeam2, in which GFP and OFP were used as a FRET pair to monitor intracellular ATP levels with an affinity $K_d$ of 2.3 mM at 37°C. Laser-based axotomy was applied (white bars in F, white arrow in G) along a distal axon. The green color is the cp173-mEGFP channel, and the red color is the OFP channel (mKO2). The ratiometric kymograph was generated by time-lapse imaging of a total of 100 frames with 5-s intervals. Axotomy was applied at the 51th frame (G). Note that axotomy triggers acute ATP depletion at millimolar levels in the vicinity of the injured site. Bars: (A–C) 10 µm; (D–G) 20 µm.
Figure 6. Enhanced mitochondrial transport recovers mitochondrial integrity and rescues energy deficits in injured axons. (A and B) Kymographs (A) and quantitative analysis (B) showing axonal mitochondrial flux of WT and snph KO cortical neurons along microgrooves. Neurons were infected with pLenti-GFP-Mito. Time-lapse imaging was recorded for 30 frames with a 5-s interval for a total of 2.5 min. In kymographs, vertical lines represent stationary organelles; oblique lines or curves to the right indicate anterograde transport toward terminals. The mitochondrial flux in axonal bundles was measured by the total events of bidirectional transport through microgrooves. The images were acquired by 40× lens. Note that deleting snph significantly increases axonal mitochondrial flux before and after axotomy when compared with WT neurons. Data were analyzed from the total number of microgrooves indicated within bars from more than three independent experiments and expressed as mean ± SE and by Student’s *t* test. (C and D) Images (C) and quantitative analysis (D) showing the recovery of TMRE staining on axonal mitochondria in the microgrooves adjacent to the axotomized site. Cortical neurons were infected

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in WT neurons by overexpressing Miro1 also recovers axotomy-induced energy deficits (Fig. 6 I). Our study thus highlights the crucial role of mitochondrial transport in maintaining ATP supply in response to axotomy-induced mitochondrial damage.

To examine whether adult *snph* KO neurons display any altered capacity of calcium buffering, we applied the green fluorescent Ca²⁺ sensor G-GECO1.2 with a $K_3 = 1.15$ µM (Zhao et al., 2011), a Ca²⁺ sensor well characterized in our previous study (Di Giovanni and Sheng, 2015). The sensor G-GECO1.2 was electroporated into neurons and imaged 2 d later. The multi-timer macro was applied for consecutive laser axotomy and image acquisition of calcium buffering capacity (Fig. S5 E). The maximal length of Ca²⁺ sensor response was unchanged after deleting *snph* ($P = 0.423$; Fig. S5 F). The mean decay time constant within 40 µm of the proximal end of injured axons, calculated based on the fluorescence intensity ($F/F_0$) curve with a nonlinear fit, showed no significant change (WT: 15.28 ± 0.08; KO: 15.57 ± 0.08, $P = 0.827$; Fig. S5 G). Thus, Ca²⁺ sensing is unlikely the key pathway in *snph* KO neurons to facilitate axonal regrowth after injury.

*snph* KO sciatric nerve displays enhanced regenerative capacity in vivo after injury.

Our ex vivo imaging in adult *snph* KO mice (2 mo old) displays enhanced mitochondrial motility along axonal bundles of sciatric nerve (71.38 ± 2.72%, $P < 0.0001$) relative to those from control littermates (30.948 ± 2.17%; Videos 1 and 2). To test whether enhanced mitochondrial transport facilitates axonal regeneration in vivo, we crushed sciatric nerve in adult *snph* KO mice and control littermates (2 mo old) and assessed the level of regeneration past the crush site 3 d after injury. We first verified SNPH depletion in *snph* KO sciatric nerve by co-immunostaining SNPH and βIII-tubulin. Although SNPH-targeted mitochondria are abundant along the axons of sciatric nerves of WT mice, SNPH staining was absent in the same region of *snph* KO mice (Fig. 7 A).

To visualize regrowth of damaged axons in vivo, longitudinval sciatric nerve sections were stained for the growth-associated protein, GAP-43. The reported concentrations of GAP-43 in growth cones (Goslin et al., 1988) and regenerating axons (Fu and Gordon, 1997) make it an ideal marker to track axon regeneration in vivo (Ackermann et al., 2002; Abe et al., 2010). GAP-43 expression was undetectable or at a very low level in the axons of uninjured WT and *snph* KO sciatric nerve (Fig. 7 B). Adult WT and *snph* KO mice were subjected to a sciatic nerve crush injury and axonal regenerative capacity 3 d after the crush injury was assessed in sciatic nerve longitudinal sections by examining GAP-43–positive axonal processes. The length of GAP-43–expressing axons past the crush site was significantly increased in *snph* KO sciatric nerves compared with that of control littermates (Fig. 7 C). We quantified this increase by normalizing the number of GAP-43–expressing axons at various distances from the crush site. *snph* KO sciatric nerves display significantly more regenerating axons at 1.5 ($P < 0.001$), 2.0 ($P < 0.01$), 2.5 ($P < 0.01$), and 3.0 mm ($P < 0.01$) distal from the crush site than WT controls (Fig. 7 D). In addition, we calculated a regeneration index by measuring the distance away from the crush site in which the average number of GAP-43–positive axons is half of that observed at the crush site. The regeneration index is significantly higher in *snph* KO sciatric nerve compared with that of WT ($P = 0.01$, Student’s $t$ test; Fig. 7 E). Thus, our in vivo crush study suggests that enhanced axonal mitochondrial transport in *snph* KO sciatric nerves accelerates axon regeneration.

Next, we examined whether rescued energy deficits in mature *snph* KO neurons impact GAP-43 expression in both injured ipsilateral and uninjured contralateral DRG neurons 3 d after injury. Although WT contralateral DRG neurons display undetectable or low levels of GAP-43, sciatric nerve injury induces its expression in ipsilateral DRG neurons (Fig. 8, A and B), consistent with injury–induced GAP-43 expression. To our surprise, GAP-43 expression in *snph* KO DRGs is increased to a larger extent after injury. Although a significant number of uninjured contralateral DRG neurons express GAP-43, which mimics the peripheral conditioning lesion effect, sciatric nerve injury further enhances its expression. Elevated GAP-43 expression is induced in association with nerve regeneration, potentially recapitulating an early developmental program. Enhanced axonal mitochondrial transport and rescued energy deficits in mature *snph* KO neurons likely activate the early intrinsic program to support neuron regrowth.

**Discussion**

In the current study, we reveal a previously unknown cellular pathway that controls axonal regenerative capacity in mature neurons. Although mitochondrial transport in axons progressively declines with neuron maturation, axonal injury triggers an acute stress signal that depolarizes local mitochondria, leading to energy deficits near the injured site. Thus, maintaining a local ATP supply via recruiting healthy mitochondria into injured axons is critical to meet the enhanced metabolic requirements during axonal regeneration. Enhancing mitochondrial transport with pBent-GFP-Mito, and axon bundles were loaded with 20 nM TMRE dye before or 1 or 5 h after axotomy. Live images were acquired at 12 bit below the saturate setting for quantitative analysis of ΔΨm, of individual GFP-Mito–labeled mitochondria in axon bundles. Note that axotomy impairs mitochondrial integrity by depolarizing ΔΨm, [reduced TMRE staining, arrowheads] in the distal axons of WT neurons; the phenotype can be effectively reversed by enhanced mitochondrial transport in *snph* KO axons. Data were analyzed from the total number of mitochondria ($n > 1000$) for each condition indicated within bars. (E and F) Pseudo-color images (E) and quantitative analysis (F) showing ATP maintenance in the growing tips 6 h after axotomy in WT and *snph* KO neurons infected with pBent-PercevalHR. The $F_{560nm}/F_{510nm}$ ratio metric mean intensity of PercevalHR reflects the relative ATP/ADP ratio. Note that enhanced mitochondrial transport in *snph* KO axons increases the ATP/ADP ratio in the growing tips (white arrows). (G and H) Pseudo-color ratiometric images (G) and the $F_{550nm}/F_{510nm}$ ratiometric integrated intensity (H) of ATP probe GO-Aleam2. Both WT and *snph* KO cortical neurons were transfected with ATP-sensitive GO-Aleam2 probe, followed by imaging distal 150-µm microgrooves before or 1, 3, or 5 h after axotomy. Note that WT neurons display reduced $F_{560nm}/F_{510nm}$, integrated intensity after axotomy, suggesting ATP depletion. Enhancing mitochondrial transport in *snph* KO neurons reverses energy deficits. (I) Overexpressing Miro1 rescues axotomy-induced energy deficits. Quantitative analysis shows the $F_{560nm}/F_{510nm}$ ratiometric integrated intensity (left) and normalized $F_{550nm}/F_{510nm}$ integrated intensity (right) of ATP probe GO-Aleam2. WT neurons were coinfected with lent-GO-Aleam2 and lent-control vector or lent-Miro1 at DIV0, followed by axotomy at DIV8 and imaging of distal 150-µm microgrooves before or 1, 3, or 5 h after axotomy. Note that enhancing mitochondrial transport in WT neurons expressing Miro1 recovers energy deficits. Data were analyzed from the total number of microgrooves indicated within bars or in parentheses and expressed as mean ± SE and Mann–Whitney U test (F) or Student’s t test (D, H, and I). Bars, 20 µm.
mitochondrial transport facilitates regeneration • Zhou et al.

not only helps remove those injury-damaged mitochondria, but also replenishes healthy ones in the injured axons, thus recovering mitochondrial integrity and reversing energy deficits. This cellular process is critical for mature neurons to facilitate their regeneration after injury (Fig. 9). Therefore, our study provides new mechanistic insights as to how mature neuron-associated elevation of SNPH expression contributes to regeneration failure. Thus, rescuing local energy deficits by enhancing mitochondrial transport is one of the potential cellular targets for new therapeutic strategies to stimulate axon regeneration.

Energy deficits in injured axons contribute to regeneration failures

CNS injury in adult mammals usually leads to a failure to initiate regeneration because of an inability to assemble a new active growth cone, a cellular process that includes sealing ruptured membranes, retrograde growth signaling, cytoskeleton restructuring, local translation, transport of building materials, and insertion of new membrane and cell surface molecules (Bradke et al., 2012). All of these steps demand high energy consumption. ATP consumption is central to axon growth and regeneration. Mitochondrial ATP production provides most of the axonal energy. Because the diffusion capacity of intracellular ATP through long axons is rather limited (Sun et al., 2013), local mitochondria provide the ATP necessary to support axonal regeneration. Axonal injury induces mitochondrial depolarization and oxidative stress associated with axonal degeneration (O’Donnell et al., 2013; Cavallucci et al., 2014). Therefore, proper mitochondrial transport into regenerating axons ensures that metabolically active areas are adequately supplied with ATP. Thus, ATP consumption and supply are central to axon regeneration.

Our study demonstrates that both physical and laser-based axotomy triggers an acute stress signal that efficiently depolarizes local mitochondria (Fig. 5 and Fig. S4), thus reducing ATP supply to the growing tips after injury (Fig. 6 and Fig. S5). Mitochondrial dysfunction and impaired transport are associated with the pathology of neurodegenerative disorders (see reviews by Chen and Chan [2009], Court and Coleman [2012], and Sheng and Cai [2012]). Dysfunc-
tional mitochondria not only supply less ATP, but also release toxic reactive oxygen species and apoptotic factors, which further trigger axonal pathology and degeneration (Lucius and Sievers, 1996; Koeberle and Ball, 1999; Alvarez et al., 2008). We show that SNPH-mediated mitochondrial anchoring in mature axons impairs delivery of healthy mitochondria to and removal of those damaged mitochondria from injured axons, thus leading to energy deficits and loss of regrowth capacity. However, the mitochondrial stress and energy deficits at the injured axons can be effectively rescued by enhanced mitochondrial transport in axons (Fig. 6). Efficiently replacing those damaged mitochondria serves as a neuroprotective mechanism essential for regeneration.

Mature neuron-associated elevation of SNPH expression inhibits regenerative capacity

The majority of axonal mitochondria are stationary in mature neurons, which is achieved through an SNPH-mediated anchoring mechanism. SNPH targets axonal mitochondria through its carboxyl-terminal mitochondria-targeting domain and immobilizes mitochondria by anchoring to MTs (Kang et al., 2008; Chen and Sheng, 2013). Deleting snph results in a robust increase of axonal mitochondrial transport. Conversely, overexpressing SNPH abolishes mitochondrial transport in axons. Strikingly, SNPH is strictly developmentally regulated in brains: its expression is hardly detectable in embryonic stages and before P7 and peaks 2 wk after birth (Das et al., 2003). Here, we further reveal mature neuron-associated SNPH expression (Fig. 2, D and E). As a result, axonal mitochondria display progressively reduced motility during maturation from DIV7 to DIV18 (Fig. 2, F and G). This unique SNPH expression highlights a new intrinsic pathway controlling axonal regrowth capacity: increased SNPH expression in mature neurons and reduced mitochondrial transport account for the mature neuron-associated decline of regenerative capacity. Therefore, the snph KO mouse is an ideal genetic model to examine the impact of enhanced mitochondrial transport on in vitro regrowth of mature neurons and in vivo regeneration in adult mice.

We found that when mitochondrial transport is abolished by genetic manipulation, axon regrowth fails after axotomy. Conversely, enhancing mitochondrial transport in mature neurons significantly increases axon regrowth in vitro and regeneration in vivo after injury (Figs. 3 and 7). Such regrowth recovery depends on mitochondrial ATP production: (a) blocking mitochondrial ATP generation abolishes the regrowth recovery in snph KO neurons, and (b) applying ATP to neurons overexpressing SNPH partially recovers regrowth capacity. Altogether, these in vitro and in vivo studies suggest that maintaining the ATP supply to injured axons by enhancing mitochondrial transport facilitates axonal regeneration.

Mitochondrial flux into distal axons supports axonal growth and branching during early brain development (Morris and Hollenbeck, 1993; Spillane et al., 2013; Tao et al., 2014). A recent study highlights a critical role for SNPH-mediated mitochondrial anchoring in the maintenance of axonal branching through the AMPK signaling pathway (Courchet et al., 2013). Activation of AMPK increases anterograde flux of mitochondria into axons and induces axonal branching in an ATP-dependent manner. As a cellular energy sensor, AMPK activation may regulate signaling pathways that replenish ATP supplies. Although these studies focused on axonal growth in developing neurons, a fundamental question remains unaddressed: does mitochondrial transport play a critical role in the facilitation of axonal regeneration in mature neurons? The snph KO neurons show no detectable changes in initial axon growth during early de-
Developmental stages when compared with WT neurons (Fig. S1, A and B; Kang et al., 2008); this is attributed to the fact that SNPH is hardly detectable during early development (Fig. 2, D and E). Thus, the SNPH-mediated mechanism controls axonal regrowth/regeneration in mature neurons, rather than initial growth in developing neurons.

Selection of in vivo system for injury-induced regeneration

The failure of CNS regeneration is caused by a lack of induction of a cell-intrinsic growth capacity after injury (Afshari et al., 2009; Giger et al., 2010) and the presence of extrinsic inhibitory effects (Fiblin, 2006; Yiu and He, 2006; Hoffman, 2010; Sun et al., 2011). In view of these, we considered that enhancing mitochondrial transport and rescuing energy deficits may not be sufficient to activate intrinsic signaling pathways and overcome extrinsic inhibitory effects in the CNS. In contrast, injured axons in the adult PNS maintain some capacity to regenerate (Abe and Cavalli, 2008; Chandran et al., 2016). We reasoned that regenerative capacity in PNS could be further facilitated in snph KO mice if enhanced mitochondrial transport rescues local energy deficits after injury. Thus, by focusing on the sciatic nerve crush study, we provide in vivo evidence showing that snph KO sciatic nerves display enhanced regenerative capacity (Fig. 7). This enhanced regeneration is consistent with robustly enhanced mitochondrial motility along snph KO sciatic nerves (71.58 ± 2.72%, P < 0.0001) relative to those from WT adult mice (30.948 ± 2.17%; Videos 1 and 2). We also found that GAP-43 expression in snph KO DRGs is increased to a larger extent after injury (Fig. 8, A and B). It is well documented that elevated GAP-43 expression is associated with nerve regeneration, potentially recapitulating an early developmental program (Fu and Gordon, 1997; Ackermann et al., 2002; Abe et al., 2010). Alternatively, enhanced mitochondrial transport in mature snph KO neurons activates the early intrinsic program to support neuron regrowth. Exploring how an enhanced energy supply in injured axons impacts the GAP-43 signaling pathway is of great interest in future research.

Our study provides mechanistic insights into how mitochondrial anchoring and energy deficits influence axonal regenerative capacity. Our model is supported by several studies manipulating mitochondrial transport in axons. In the Caenorhabditis elegans mutant ric-7, in which mitochondria fail to transport to distal axons, injured axons degenerate rapidly; such degeneration can be suppressed by forcing mitochondria into the axons (Rawson et al., 2014). In an adult fly model, Fang et al. (2012) revealed that axonal mitochondria are rapidly depleted upon axotomy. The injury-induced mitochondrial loss is suppressed by up-regulation of Nmnat, which is known to suppress axon degeneration (Gilley and Coleman, 2010). Interestingly, when mitochondria are genetically eliminated from axons by depleting the kinesin adaptor Milton, up-regulation of Nmnat is no longer effective to suppress axon degeneration (Fang et al., 2012). Similarly, WldS (slow Wallerian degeneration) is an effective protein that suppresses Wallerian degeneration after axon injury (Conforti et al., 2000; Coleman and Freeman, 2010). Using both Drosophila melanogaster and mouse models, Avery et al. (2012) identified axonal mitochondria as a key site of action for WldS neuroprotective function. Altogether, these studies consistently suggest that regulating mitochondrial transport is a common target for the signaling pathways protecting injured axons from degeneration.

In summary, our study addresses three fundamental issues as to whether (a) axonal mitochondrial transport is essential for mature neurons to regenerate, (b) enhancing mitochondrial transport facilitates axonal regenerative capacity both in vitro and in vivo, and (c) enhanced mitochondrial transport rescues local energy deficits in injured axons. Our study highlights that reduced mitochondrial transport in mature neurons and injury-induced energy deficits reflect common factors limiting regenerative ca-
pacity. Our in vivo and in vitro experiments suggest that activating an intrinsic “growth program” requires the coordinated modulation of mitochondrial transport and recovery of energy deficits. Such combined approaches may represent a valid therapeutic strategy to facilitate regeneration in the injured CNS and PNS.

Materials and methods

Mouse lines and animal care

The snph<sup>−/−</sup> mouse line was generated by targeted gene replacement in embryonic stem cells as previously described (Kang et al., 2008). Thy1/Mito-CFP transgenic mice were purchased from the Jackson Laboratory. The mice and the Sprague-Dawley rats were maintained in the National Institute of Neurological Disorders and Stroke (NINDS) animal facility and housed in a 12-h light/dark cycle. All animal procedures were performed according to National Institutes of Health (NIH) guidelines and were approved by the NINDS/National Institute on Deafness and Other Communication Disorders (NIDCD) Animal Care and Use Committee.

Antibodies

The purified polyclonal antibody against SNPH residues 225–428 was described previously (Kang et al., 2008). Sources of other antibodies or reagents are as follows: anti-βIII-tubulin, anti–MAP2, anti–Tau-1, anti-dynein IC74, and anti–GAP-43 were from EMD Millipore; anti-cytochrome c was from BD; anti–TOM20 was from Santa Cruz Biotechnology, Inc.; anti-Miro1/2 (HPA010687) and anti-Trak2 were from Sigma-Aldrich; ECL-HRP–linked secondary antibodies were from GE Healthcare; and Alexa Fluor 546– or Alexa Fluor 488–conjugated secondary antibodies were from Invitrogen.

Lentivirus infection

The cDNA gene encoding SNPH, SNPH-dMTB, or Miro1 was cloned into the lentivirus vector pHUGW with the CMV promoter and HA tag; the YFP, GFP-Mito, or PercevalHR (Addgene) and GO-ATeam2 and GO-ATeam3 (gifts from H. Imamura, Kyoto University, Kyoto, Japan) were cloned into pHUGW vector. All lentivectors were prepared and transduced at the same concentration. For infection, 2 × 10⁵ freshly dissociated neurons were incubated with up to 5 µl concentrated virus preparation in 25 µl culture medium for 1 h; 200 µl culture medium was then added to each side of the wells.

Microfluidic neuron culture and axotomy

For axotomy and staining, both commercialized (#RD450; Xona) and house-made microfluidic devices were used. For the latter, a silicon wafer with a pattern made out of SU-8 by photolithography was used to cast the PDMS microfluidic devices. In brief, SYL Gardner 184 silicone elastomer base was mixed with the curing agent at a ratio of 10:1. The PDMS was then mixed well using a THINXY mixer ARF-310 in two steps: mixing at 2,000 rpm for 4 min and de-foaming at 2,200 rpm for 4 min. The well-mixed PDMS was poured onto the silicon wafer and then placed in a Bel-Art vacuum desiccator for 3 h to help remove air bubbles from the PDMS. The wafer with PDMS was placed in an 80°C oven for 1–2 h to cure. Once the PDMS was cured, the PDMS was pulled out, and reservoirs were punched out. The PDMS devices were extensively washed and autoclaved before use. The coverslips and house-made devices were exposed to the plasma treatment for 2 min in a PDC-320 plasma cleaner and bonded together for neuronal culture.

Brain tissues were dissected out from E18–19 mouse embryos and kept in ice-cold Hibernate buffer supplemented with 2% B27 and antibiotics. After the embryos were genotyped by direct tail PCR, cortical neuron cultures were prepared with papain as described previously (Kang et al., 2008). After preparing a single-cell suspension, 2 × 10⁷/20 µl dissociated neurons were added into the cell body chamber of the microfluidic devices. After cells were attached, 100 µl of prebalanced culture medium was added into each well of the device. Axons pass through microgrooves into the axonal terminal chamber at DIV4 as they grow. For axon regrowth assays, axons in terminal chambers were axotomized by vacuum aspiration at DIV5–7 or DIV12. The two wells on the axon sides were subjected twice to aspiration to ensure complete axotomy. After the first aspiration, Hibernate medium was added to the axon chambers, followed by the second aspiration. The devices were then refilled with fresh culture medium for further incubation for various times, as indicated.

For quantitative analysis of axon growth before and after axotomy, axons in the terminal chamber labeled by lentivirus or βIII-tubulin staining, were imaged under the same setting using an 880 confocal microscope (ZEISS) with a 20× objective. The area of YFP or βIII-tubulin signals above the same threshold within a 1024 × 1024 image that covers all axon segments extending from microgrooves was measured using ImageJ (NIH). Images were collected from multiple chambers in at least three experiments for quantitative analysis.

Adult DRG neuron cultures and Sholl analysis

Adult WT and SNPH KO littermates at P60 were dissected in HBSS. After clipping off bones from the spinal cords, DRGs were pulled out and the excess roots were trimmed off in the HBSS buffer. DRGs were digested in 2.5 U/ml dispase II (Roche) and 200 U/ml collagenase (Worthington Biochemical) for 30 min at 37°C and then an additional 35 min at 30°C. DRG neurons were collected and triturated in Neurobasal-A medium supplemented with B27 and 0.5 mM GlutaMAX (Invitrogen). After purification with 1% BSA cushions, the isolated DRG neuronal cell bodies were plated onto coverslips coated with 30 µg/ml poly-L-ornithine and 5 µg/ml laminin (Roche) at a final density of 2,000 cells per 12-mm coverslip. For axotomy, one-d culture in vitro, DRG neurons were stained for imaging, and a Sholl analysis of total numbers of axon intersections was performed with an ImageJ plugin.

Time-lapse imaging and kymograph analysis

To image mitochondrial transport in axons, neurons grown on the PDMS device were replaced with prewarmed Hibernate A low fluorescence medium (BrainBits) supplemented with 2% B27, 0.5 mM GlutaMAX, and 1% Pen/Strep at day of image. The temperature was maintained at 37°C during time-lapse imaging. The microgrooves of the device that contain the fasciculated axon bundles were visualized. Time-lapse imaging of the microgrooves was recorded using a confocal microscope (LSM 510 META; ZEISS) with a C-Apochromat 40×/1.3 oil NA objective (ZEISS). The image sequences were collected at a 512 × 512–pixel resolution (12 bit) with 5-s intervals at a scanning zoom of 1.0 in the x and y directions; 100–200 frames were captured for each condition. For mitochondrial transport before and after axotomy, only 30 frames were recorded to coherently match the time window. To quantify motility, a kymograph was generated by ImageJ plugin as previously described (Kang et al., 2008). For each experimental group, all visible vesicles on the kymograph were pooled together. The mitochondria were considered stationary if they remained immobile during the entire recording period; a mobile one was counted if the net displacement was >5 μm.

Microscope image acquisition

For imaging acquisition, a prescan of all samples was conducted to ensure confocal settings below saturation at 1024 × 1024 pixels (12 bit). From a given experiment, all images were acquired on the same
day with new pinhole adjustment and under the same confocal settings. Morphometric measurements were performed using ImageJ. For ATP/ADP ratio analysis, the neurons transfected with PercevalHR were excited using 488/458 filters, and the emissions were collected at 505–550 nm. For axonal ATP analysis, two emission images were collected at 505–550 nm and long pass above 545 nm along the GO-A-Team2 or GO-A-Team3 expression axon. Radiometric images were generated by an ImageJ macro using binary Mask image to remove nonspecific background in the microgrooves. To analyze axon regrowth, a fixed threshold was manually set to ensure that most signals were detected but not oversaturated with a minimum detectable background. Both the pixel area and fluorescence integrated intensity were measured. For active growth cone quantification, the axon tips that were labeled with 543 phalloidin and βIII-tubulin were imaged and a fixed threshold and pixel size were applied for ImageJ analysis. Other parameters are described in the figure legends.

Measurement of mitochondria membrane potential (ΔΨm)
Lenti-GFP-Mito was introduced into cultured neurons to label total mitochondria. Before and right after axotomy, nonplasma-treated devices were removed and the exposed axon bundles along microgrooves were loaded with 25 mM TMRE (Invitrogen) for 30 min and then washed three times with B27/Hibner A low fluorescence medium before imaging. ΔΨm was assessed by relative TMRE fluorescent intensity of each individual mitochondrion labeled by GFP-Mito along the most distal 200 μm of axon bundles on microgrooves using ImageJ particle analysis.

ATP rescue experiments
Neurons were electroporated by Nucleofector (Amaxa) using program O-004 with an electroporation buffer (86 mM KH2PO4, 17.5 mM NaHCO3, 2.72 mM d-glucose, and 11.6 mM MgCl2-6H2O) containing 7.1 mM ATP disodium, followed by spin down and removal of the supernatant containing ATP. For ATP rescue study, the electroporated neurons were immediately plated onto a microfluidic chamber with medium containing the final ATP concentration adjusted to 200 μM.

Acquisition of axotomy time-lapse imaging
The multi-timer macro was applied for a combination of laser-based axotomy and time-lapse imaging, as previously described (Yanik et al., 2004; Cho et al., 2013) with some modifications. For axotomy, an LSM 510 META two-photon confocal microscope (ZEISS) with an inverted P-Apochromat 40×/1.3 oil objective was applied. After alignment of scan head and collimator adjustment, Mira 900F laser power was set on to initiate mode-lock without CW break. The stable mode-lock was achieved by tuning the slits on the Coherent Mira 900F. Once the arbitrary units in the power indicator were above 260, the beam wavelength was set to 730 nm for bleaching. For consecutive real-time imaging capture of TMRE and GFP-Mito before and after axotomy, both acquisition and bleach parameters were preset and reloaded in multi-timer macro blocks. In brief, the images in the first block were first recorded at 5-s intervals for a total of 50 frames by 400/543-nm laser in line switch mode; then, the second block reloaded the 730-nm laser that powered-up with 70% output and pixel dwell time >10 μs in cropped imaging region for bleaching. The post-axotomy recording was collected at 5-s intervals for a total of 50 or 100 frames by 488/543-nm laser in the final block.

To measure the Ca2+ buffering capacity in the adult DRG axons, the neurons were electroporated with green fluorescent Ca2+ sensor G-GECO1.2. The same multi-timer macro was applied for consecutive laser axotomy and image acquiring. In brief, after 2 frames of base line acquisition and two-photo laser, 730 nm was applied at the third frame, and then another 26 frames were collected at 512 × 512–pixel resolution (12 bit) with 3-s intervals. The average decay time constant was calculated based on fluorescence intensity (F/F0) curve with nonlinear fit.

For physical axotomy and time-lapse imaging, the pure axon bundle was first stained by TMRE and MitoTracker Green, and images were acquired before and after physical axotomy by using pulled glass capillaries (World Precision Instruments) with a tip diameter of ~1 μm that were prepared from Sutter Instrument (p-97).

In vivo axonal regeneration after injury
For in vivo sciatic nerve injury and axonal regeneration, adult snph KO and WT littermates (2 mo old) were subjected to unilateral sciatic nerve crush as described previously (Abe et al., 2010). In brief, under deep anesthesia, the left sciatic nerve was exposed at the mid-thigh level; the nerve was then crushed using fine hemostatic forceps three times, 15 s each time. The wound was sutured in layers, and the mice were allowed to recover on a heating pad. 3 d after injury, mice were perfused transcardially with PBS (pH 7.4) followed by ice-cold 4% PFA in PBS. Sciatic nerves and L4/L5 DRGs from the injured side as well as the contralateral uninjured side were dissected, post-fixed in 4% PFA in PBS for 2 h, and then cryoprotected overnight in 30% sucrose. Specimens were embedded in tissue-freezing medium and then cut longitudinally into serial sections at a thickness of 12 μm. Immunohistochemistry was performed to evaluate axonal regeneration with a GAP-43 antibody to label regenerating axons. Sciatic nerve sections were first blocked with 0.3% normal goat serum in PBS containing 0.2% Triton X-100 and then incubated with primary antibodies for 1 h at room temperature. The following primary antibodies were used: rabbit anti-GAP-43 antibody (1:800; Abcam), mouse anti-βIII-tubulin antibody (1:1,000; Sigma-Aldrich), or rabbit anti-SNPH antibody (1:400). Sections were washed with PBS, followed by incubation with Alexa Flour 546–conjugated goat anti–rabbit IgG or Alexa Flour 488–conjugated goat anti–mouse IgG (1:500; Invitrogen) secondary antibodies together with DAPI (1:1,000) to visualize nuclei. Slices were mounted and images were acquired using an LSM 880 confocal microscope (ZEISS) with a 20× objective. Three sections spaced 60 μm apart for each sciatic nerve were used to assess axonal regeneration. The nerve was divided into 0.5-mm segments; the number of axons in the segment 0.5 mm proximal to the injury site (0 mm) and in the distal 0.5-, 1.0-, 1.5-, 2.0-, 2.5-, 3.0-, and 3.5-mm segments were quantified and normalized to the number in the injury site.

To examine GAP-43 expression in DRG neuronal cell bodies in response to sciatic nerve injury, 9-wk-old snph KO and their WT littermates were subjected to unilateral sciatic nerve crush injury. DRG sections (three sections each) from ipsilateral and contralateral sides of both WT and snph KO mice were co-immunostained with antibodies against GAP-43, βIII-tubulin, and DAPI 3 d after sciatic nerve crush injury. Images were obtained using an LSM 880 confocal microscope with a 40× objective. The mean intensity of GAP-43 in individual DRG neuronal cell bodies was measured using ImageJ and normalized to the mean intensity of that from the contralateral uninjured side of WT mice.

Statistical analysis
Statistical analysis was performed using Prism (GraphPad Software). Two groups were compared using F-test, Mann–Whitney (sample size \( n < 30 \)), or Student’s t test (sample size \( n \geq 30 \)). For multiple comparisons, one-way ANOVA analysis (\( n \geq 30 \)) or Kruskal–Wallis test (\( n < 30 \)) is used. Data are expressed as mean ± SEM. Differences were considered significant with \( P < 0.05 \).

Online supplemental material
Fig. S1 shows that deleting snph did not change early axonal growth and Miro1/2 expression is not regulated by SNPH. Fig. S2 shows...
that enhanced axonal regrowth in snph KO neurons is not directly correlated with cell density in the somatic chambers. Fig. S3 shows that overexpressing SNPH alters mitochondrial distribution and ATP/ADP ratios in distal axons. Fig. S4 shows that axotomy stress induces mitochondrial depolarization. Fig. S5 shows energy deficit recovery and calcium buffering in snph KO neurons after axotomy. Videos 1 and 2 show ex vivo imaging of axonal mitochondrial transport along sciatic nerves in Thy1-Mito-CFP and in the crossed snph KO/Thy1-Mito-CFP mice, respectively. Videos 3–5 show axonal mitochondrial motility along microgrooves from cortical neurons overexpressing SNPH, SNPH-dMTB, or Miro1, respectively. Videos 6 and 7 show axotomy stress depolarizing mitochondria in the vicinity of injured sites. Video 8 shows axonal mitochondrial motility along microgrooves from WT cortical neurons 1 h after axotomy. Video 9 shows enhanced mitochondrial motility along microgrooves from snph KO cortical neurons 1 h after axotomy. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201605101/DC1.

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Animal care and use were carried out in accordance with NIH guidelines and approved by the NIH, NINDS/NINDCD Animal Care and Use Committee.

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The authors declare no competing financial interests.

Author contributions: B. Zhou designed and conducted in vitro and in vivo experiments and data analysis; P. Yu designed in vivo experiments and data analysis; M.-Y. Lin performed biochemical analysis; Y. Chen characterized ex vivo mitochondrial transport; T. Sun performed ATP/ADP assays with pH calibration; Z.-H. Sheng is the senior author who conceived and designed the project; B. Zhou and Z.-H. Sheng wrote the manuscript.

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References


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Figure S1. **Deleting snph does not change early axonal growth, and Miro1/2 expression is not regulated by SNPH.** (A and B) Representative images (A) and quantitative analysis (B) showing no observable change in early axonal growth of snph KO neurons before axotomy. WT and snph KO cortical neurons grown on microfluidic chambers were immunostained with βIII-tubulin at DIV5. Note that the normalized axonal area grown from each microgroove was not significantly altered between WT and snph KO neurons (P = 0.38). (C) Representative time-lapse images of live neurons showing regenerating axons from the axotomized tips. The snph KO cortical neurons were infected with lentivirus and axotomized in axonal terminal chambers at DIV10. Time-lapse images were captured at 0, 4, 6, and 8 h after axotomy. White and yellow arrows point to axotomized sites. Data were collected from the total number of microgrooves shown in the bars and expressed as mean ± SE and statistically analyzed by paired Student’s t test. Bars: (A) 100 µm; (C) 200 µm; (D and E) Representative immunoblots and quantitative analysis showing that cortical neurons (DIV14) deleting snph (D) or overexpressing SNPH (E) display no detectable change in Miro1/2 expression. WT cortical neurons were infected by lentivirus HA-SNPH or HA control, and then Miro1/2 expression was examined at DIV14. Equal amounts (20 µg) of cell lysates from cultured neurons were loaded and sequentially immunoblotted with antibodies on the same membranes after stripping between applications of each antibody. Note that the snph KO or WT neurons overexpressing HA-SNPH do not change Miro1/2 expression. The relative intensity of Miro1/2 was quantified from three repeats and normalized to actin. Data are expressed as mean ± SE and statistically analyzed by paired Student’s t test.
Figure S2. Enhanced axonal regrowth in snph KO neurons is not directly correlated with cell density in the somatic chambers. (A–C) Representative images (A) and quantitative analysis showing normalized MAP2-positive neurons (B) and the percentage of TUNEL-positive death cells (C) in the somatic chambers. 0.2 x 10^6 [10^7/ml] E18 mouse cortical neurons were plated in the somatic chamber, and axotomy was performed at DIV7 in the axonal terminal chamber. The number of MAP2-labeled neurons in the somatic chamber was counted before and 1 h and 3 d after axotomy, and data were normalized to that from neurons before axotomy. The percentage of TUNEL-positive cells to total DAPI staining in each somatic image field was calculated based on automatic watershed binary images. Note that no significant change was observed in the normalized density of MAP2-positive neurons and the percentage of TUNEL-positive death cells between WT and snph KO neurons at 1 h and 3 d after axotomy (P > 0.05). (D) Relative correlation between neuron density and axonal regrowth after injury. Although WT soma density correlates with axonal regrowth after injury (R^2 = 0.3501, P < 0.0001), enhanced axonal regrowth in snph KO neurons is not directly correlated with relative neuron density in the somatic chamber (R^2 = 0.0556, P = 0.1187). Data were collected from the total number of images (425 x 425 µm^2) shown in the bars and are expressed as mean ± SE and analyzed by one-way ANOVA, Kruskal–Wallis test.
Figure S3. Overexpressing SNPH alters mitochondrial distribution and ATP/ADP ratio in distal axons. (A) Quantitative analysis showing relative mitochondrial distribution along an entire axon of cortical neurons coexpressing DsRed-Mito with GFP or together with SNPH, SNPH-dMTB, or Miro1. Neurons at DIV5 were immunostained with Tau. (B) Quantitative analysis showing relative cellular ATP/ADP ratio along an entire axon of cortical neurons coexpressing DsRed-Mito with GFP or SNPH. The cellular ATP/ADP ratio was measured by applying an engineered fluorescent ATP sensor PercevalHR in live neurons. The fluorescence intensity ratio (F488nm/F405nm) of Perceval reflects the relative ATP/ADP ratio. Relative mitochondrial distribution and ATP/ADP ratios along an axon were analyzed based on five equally divided segments (20%) of each individual axon from the most proximal to the most distal region. Note that overexpressing SNPH in cortical neurons redistributes mitochondria to the proximal region (P < 0.001), thus reducing mitochondrial density and ATP/ADP ratio in the most distal axon segment (P < 0.001 and P < 0.05, respectively). (C and D) Measurement of axonal pH using the pH dye SNA RF-5F. Axonal pH was analyzed based on 20 equally divided segments (5%) of each axon from the most proximal to the most distal region. SNA RF-5F was excited at 488 nm and detected around 580 and 640 nm. The ratio (640/580 nm; C) and calibrated pH value (D) were calculated at each 5% segment of axons as indicated. Calibrated pH values were averaged from a total of six axons. Note that pH values in an entire axon of both WT and SNPH-overexpressing neurons undergo very minor changes (7.10–7.15). Thus, reduction in the ATP/ADP ratios in distal axons expressing SNPH is physiologically relevant to reduced axonal mitochondrial density. (E and F) SNARF-5F signal was calibrated using various buffered solutions containing high K+/nigericin with varying pH values. Axons were imaged in Tyrode’s solution for 15 min, followed by a 30-min incubation in high K+/nigericin, pH 7.5, solution. Then the SNA RF-5F signal was calibrated by varying pH solutions. Images were taken once per minute. Data were analyzed from the total number of axons indicated in parentheses from three experiments and expressed as mean ± SE and by one-way ANOVA test.
Figure S4. Axotomy stress induces mitochondrial depolarization. (A) Representative images showing physical axotomy-induced mitochondrial depolarization. A physical axotomy was applied in the axonal chamber by using pulled glass capillaries (World Precision Instruments) with a tip diameter of ∼1 µm that were prepared from Sutter Instrument (p-97). Neurons at DIV10 were loaded with 20 nM TMRE (Δψm-dependent dye) and 10 nM MitoTracker Green FM (Δψm-independent dye) before axotomy. The same image field was taken showing mitochondrial Δψm before and after physical axotomy. Note that after a physical axotomy, TMRE staining of mitochondria is lost near the axotomized ends, whereas MitoTracker Green FM staining remains. (B) Representative images showing that laser axotomy does not reduce the signal intensity of MitoTracker Red FM. The axonal chambers were loaded with 10 nM MitoTracker Green FM and 20 nM MitoTracker Red FM (Δψm-insensitive dye with similar wavelength as TMRE). A 730-nm laser-induced axotomy was applied in the axonal chamber. The same image field was taken showing mitochondria at 5 s before and 15 s after laser axotomy. Note that laser axotomy does not reduce signal intensity of MitoTracker Red FM, thus excluding an artificial effect of 730-nm laser on TMRE signals with the similar wavelength. (C) Quantitative analysis showing that somatic mitochondria maintain Δψm after axotomy. Somatic mitochondrial integrity was examined by loading MitoTracker Orange-CMTMRos, a fixable Δψm-dependent dye, before and 1 and 5 h after injury in WT and KO cortical neurons. Note that there is a subtle increase in the mean intensity of Orange-CMTMRos at 1 h after injury in both WT and KO neurons. Data were collected from the total number of images (425 × 425 µm²) shown in the bars and are expressed as mean ± SE and analyzed by one-way ANOVA, Kruskal–Wallis test. Bars, 100 µm.
Figure S5. Energy deficit recovery and calcium buffering in snph KO neurons after axotomy. (A) Measurement of pH at regrowing axonal tips using the pH dye SNARF-5F. SNARF-5F was excited at 488 nm and detected at 580 and 640 nm. The ratio (640/580 nm) was calculated from regrowing axons 6 h after axotomy, and relative pH was calibrated on SNARF-5F signal with various buffered solutions containing high K/nigericin with varying pH values (also see Fig. S3 E). Note that axon terminal pH values in both WT and snph KO neurons undergo very minor changes (7.10–7.15). Data were analyzed from the total number of axon terminals indicated in parentheses from three experiments and expressed as mean ± SE and by one-way ANOVA test. (B) Normalized F560nm/F510nm ratiometric integrated intensity of ATP probe GO-ATeam2 and ATP-insensitive GO-ATeam3 mutant (R122K/R126K). Both WT and snph KO cortical neurons were transfected with ATP-sensitive GO-ATeam2 probe or ATP-insensitive GO-ATeam3 mutant, followed by imaging along microgrooves before or 1, 3, or 5 h after axotomy. F560nm/F510nm ratiometric integrated intensity was normalized to the ratiometric intensity before axotomy. Data were analyzed from the total number of microgrooves indicated within bars and expressed as mean ± SE and Student’s t test. (C and D) Pseudo-color ratiometric images (C) and the F560nm/F510nm ratiometric intensity (D) of ATP-insensitive GO-ATeam3 mutant (R122K/R126K). Both WT and snph KO cortical neurons were transfected with ATP-insensitive GO-ATeam3 mutant, followed by imaging distal 150-µm microgrooves before or 1, 3, or 5 h after axotomy. Note that WT neurons expressing ATP-insensitive GO-ATeam3 mutant do not show a significant change in the F560nm/F510nm ratiometric intensity after axotomy. Data were analyzed from the total number of microgrooves indicated within bars and expressed as mean ± SE and Student’s t test. Bars, 20 µm. (E–G) Representative images (E) and quantitative analysis (F and G) showing similar axotomy-induced Ca²⁺ waveform between adult WT and KO DRG neurons. The green fluorescent Ca²⁺ sensor G-GECO1.2 (Kd = 1.15 µM; Di Giovanni and Sheng, 2015) was electroporated into adult DRG neurons at DIV0 and imaged 2 d later. The multi-timer macro was applied for consecutive laser axotomy and image acquiring (E). In brief, after two frames of baseline acquisition, two-photon laser 730 was applied at the third frame, and then another 26 frames were collected at 512 × 512-pixel resolution (12 bit) with 3-s intervals. The white dotted lines (E) or black bar (G) indicates laser-induced axotomy. Note that the maximal distance of Ca²⁺ sensor response was unchanged after deleting snph (P = 0.423; F), and the mean decay time constant within 40 µm of the proximal end of injured axons (black bars under images), calculated based on fluorescent intensity (F/Fo) curve with a nonlinear fit, was not significantly changed [WT: 15.28 ± 0.08, KO: 15.57 ± 0.08, P = 0.827; G]. Data are expressed as mean ± SE and were statistically analyzed by Mann–Whitney U test.
Video 1. **Ex vivo imaging of axonal mitochondrial transport along the sciatic nerve in Thy1-Mito-CFP mice.** Mitochondrial transport was assessed in sciatic nerve explants after rapid dissection and immediate immersion in prewarmed oxygenated Neurobasal-A medium. Time-lapse images of multiple axons in nerve explants were acquired using an LSM 510 META confocal microscope (ZEISS) with a P-Apochromat 40×/1.3 oil objective at 3-s intervals for a total of 100 frames. Image stacks were processed using the free ImageJ software v1.43 and associated plugins.

Video 2. **Ex vivo imaging of axonal mitochondrial transport along the sciatic nerve in the crossed snph KO/Thy1-Mito-CFP mice.** Thy1-Mito-CFP transgenic mice were crossed with C57BL/6 background snph KO/− mice to generate snph KO/Mito-CFP mice. Mitochondrial transport was assessed in sciatic nerve explants from the crossed snph KO/Mito-CFP mice. The sciatic nerve was dissected rapidly and immediately immersed in prewarmed oxygenated Neurobasal-A medium. Time-lapse images of multiple axons in nerve explants were acquired using an LSM 510 META confocal microscope (ZEISS) with a P-Apochromat 40×/1.3 oil objective at 3-s intervals for a total of 100 frames. Image stacks were processed using the free ImageJ software v1.43 and associated plugins. Note that deleting snph robustly enhances axonal mitochondrial transport along axonal bundles of sciatic nerve.

Video 3. **Axonal mitochondrial motility along microgrooves from cortical neurons overexpressing SNPH.** Cortical neurons were infected with lentivirus coexpressing DsRed-Mito and SNPH. Isolated axon bundles in the microgrooves were selected for time-lapse imaging at DIV12. Time-lapse sequences were collected at 5-s intervals for a total of 100 frames. Note that expressing SNPH abolishes mitochondrial transport in axons. Left side is toward the somatic chamber.

Video 4. **Axonal mitochondrial motility along microgrooves from cortical neurons overexpressing SNPH-dMTB.** Cortical neurons were infected with lentivirus coexpressing DsRed-Mito and SNPH-dMTB. Isolated axon bundles in the microgrooves were selected for time-lapse imaging at DIV12. Time-lapse sequences were collected at 5-s intervals for a total of 100 frames. Left side is toward the somatic chamber.

Video 5. **Axonal mitochondrial motility along microgrooves from cortical neurons overexpressing Miro1.** Cortical neurons were infected with lentivirus coexpressing DsRed-Mito and Miro1. Isolated axon bundles in the microgrooves were selected for time-lapse imaging at DIV12. Time-lapse sequences were collected at 5-s intervals for a total of 100 frames. Note that expressing Miro1 enhances mitochondrial transport in axons. Left side is toward the somatic chamber.
Video 6. Axotomy stress depolarizes mitochondria in the vicinity of injured sites. Mature cortical neurons grown on microfluidic chambers were infected with pLenti-GFP. Axons in the terminal chambers were loaded with 25 nM TMRE dye at DIV12, followed by laser axotomy and time-lapse imaging. In brief, for real-time image capture of laser axotomy, both acquisition and bleach setting were reused in separate blocks. The images were first recorded at 5-s intervals for a total of 50 frames, and then the 730-nm laser was powered-up with 70% output and pixel dwell time >20 µs in cropped imaging region for bleaching. The consecutive post-axotomy recording was collected at 5-s intervals for a total of 100 frames. For a successful laser axotomy, the axons in the DIC image were observed with a structural break down or invasion of extracellular dye. Note that GFP-labeled axons are quickly broken up upon axotomy (left) and mitochondria in the vicinity suddenly lose their TMRE staining (right).

Video 7. Axotomy stress depolarizes mitochondria in the vicinity of injured sites. Neurons were infected with pLenti-GFP-Mito, and axon bundles in the terminal chambers were loaded with 25 nM TMRE dye at DIV12. Laser axotomy and time-lapse imaging were used to acquire axon bundles with a 40x lens. The images were first recorded at 5-s intervals for a total of 50 frames; then the 730-nm laser was powered-up with 70% output and pixel dwell time >20 µs in the cropped imaging region for bleaching. The consecutive post-axotomy recording was collected at 5-s intervals for a total of 50 frames. Note that axotomy triggers a sudden loss of mitochondria staining by TMRE near the injured site, whereas those depolarized mitochondria maintain their GFP-Mito signals, and thus the color of mitochondria was switched from yellow (healthy) to green (depolarized).

Video 8. Axonal mitochondrial motility along microgrooves from WT cortical neurons 1 h after axotomy. Neurons were infected with pLenti-GFP-Mito. Time-lapse images were acquired in isolated axon bundles with a 40x lens and recorded for a total of 100 frames at 5-s intervals. Left side is toward the somatic chamber.

Video 9. Enhanced mitochondrial motility along microgrooves from snph KO cortical neurons 1 h after axotomy. Neurons were infected with pLenti-GFP-Mito. Time-lapse images were acquired in isolated axon bundles with a 40x lens for a total of 100 frames at 5-s intervals. Note that deleting snph significantly enhances axonal mitochondrial transport. Left side is toward the somatic chamber.

Reference