

Novel Tools and Methods

An ATF3-CreERT2 Knock-In Mouse for Axotomy-Induced Genetic Editing: Proof of Principle

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Abstract

Genome editing techniques have facilitated significant advances in our understanding of fundamental biological processes, and the Cre-Lox system has been instrumental in these achievements. Driving Cre expression specifically in injured neurons has not been previously possible: we sought to address this limitation in mice using a Cre-ERT2 construct driven by a reliable indicator of axotomy, activating transcription factor 3 (ATF3). When crossed with reporter mice, a significant amount of recombination was achieved (without tamoxifen treatment) in peripherally-projecting sensory, sympathetic, and motoneurons after peripheral nerve crush in hemizygotes (65-80% by 16 d) and was absent in uninjured neurons. Importantly, injury-induced recombination did not occur in Schwann cells distal to the injury, and with a knock-out-validated antibody we verified an absence of ATF3 expression. Functional recovery following sciatic nerve crush in ATF3-deficient mice (both hemizygotes and homozygotes) was delayed, indicating previously unreported haploinsufficiency. In a proof-of-principle experiment, we crossed the ATF3-CreERT2 line with a floxed phosphatase and tensin homolog (PTEN) line and show significantly improved axonal regeneration, as well as more complete recovery of neuromuscular function. We also demonstrate the utility of the ATF3-CreERT2 hemizygous line by characterizing recombination after lateral spinal hemisection (C8/T1), which identified specific populations of ascending spinal cord neurons (including putative spinothalamic and spinocerebellar) and descending supraspinal neurons (rubrospinal, vestibulospinal, reticulospinal and hypothalamic). We anticipate these mice will be valuable in distinguishing axotomized from uninjured neurons of several different classes (e.g., via reporter expression), and in probing the function of any number of genes as they relate to neuronal injury and regeneration.

Key words: Activating Transcription Factor 3; gene editing; peripheral nerve; Schwann cells; sensory neurons; spinal cord injury

Significance Statement

Understanding reactions to neurotrauma and overcoming obstacles to neural regeneration should benefit from the ability to genetically label or otherwise edit the genome of injured neurons. We sought to achieve this in mice by driving Cre recombinase expression under the control of activating transcription factor 3 (ATF3), which is robustly induced by axotomy in several populations of peripheral and central neurons. When crossed with reporter mice, recombination occurred only in injured neurons following sciatic nerve injury or spinal hemisection. Peripheral nerve injury (PNI)-induced neuronal phosphatase and tensin homolog (PTEN) excision also resulted in improved regeneration and more complete functional recovery. These results demonstrate the feasibility and utility of axotomy-induced recombination and represent a new tool for investigating genetic control of injury responses and regeneration.



Introduction

Advances in genome editing techniques have created opportunities to dissect out the function of specific genes and their contributions to health and disease. Among the most widely used of these tools is the Cre-Lox system, in which Cre expression can be restricted spatially, to a single tissue or population of cells through insertion under a specific promoter (Wagner et al., 1997; Clausen et al., 1999; Agarwal et al., 2004), or temporally, by fusing the protein to a mutated estrogen receptor selective for tamoxifen (Indra et al., 1999; Hayashi and McMahon 2002; Leone et al., 2003). In neurotrauma, the combination of spatial and temporal control of gene expression has facilitated the identification of the function of individual neural cells (Sainsbury et al., 2002; Q. Liu et al., 2010), and manipulated their phenotype to promote repair of the damaged nervous system (Park et al., 2008; Sun et al., 2011). A limitation in applying Cre-Lox to investigations of neurotrauma is the inability to selectively express Cre in populations of neurons affected by unique pathologies, such as tau protein-associated degeneration, immunemediated degradation, or simple axotomy. This could notionally be achieved by taking advantage of a gene that is highly expressed only when the neuron is degenerating or axotomized. Pathology-induced recombination would be of use to both identify the affected cell populations and to edit those populations' genetic makeup. Here, we present a novel transgenic approach to effecting genome editing on axotomy via an injury-specific gene.

Activating transcription factor 3 (ATF3) is a transcription factor that belongs to the basic leucine zipper family (Liang et al., 1996). ATF3 is an immediate early gene; its transcription is initiated extremely rapidly following the appropriate stimulus. ATF3 is considered a reliable marker of neuronal somata with injured peripheral axons (Tsujino et al., 2000; Mason et al., 2003); ATF3 mRNA can be detected after peripheral axotomy as early as 6 h after injury (Tsujino et al., 2000) and can achieve a 130-fold increase 72 h after axotomy (Seijffers et al., 2007). Other genes upregulated following injury include c-Jun and GAP-43 (Tetzlaff et al., 1991; Broude et al., 1997), but none are upregulated as fast and prominently as ATF3, making it the most reliable marker of peripheral nervous system (PNS) injury. These changes in gene expression constitute part of the pro-regenerative "cell body response" to injury. A loss of ATF3 function has been shown

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to reduce the regeneration observed after a peripheral nerve injury (PNI; Gey et al., 2016).

Damage to the PNS is a common outcome of motor vehicle accidents, penetrating trauma, and falls (Kouyoumdjian, 2006). The PNS has the ability to regenerate injured axons that result in full functional recovery: however, despite the theoretical regenerative capacity of the PNS, clinical PNI often results in permanent disability, since regeneration is often incomplete when the distance from the axotomy to the soma is far, the gap between distal and proximal stumps is large, or the time until surgical intervention is long. Reports suggest that $\sim\!5\%$ of all patients admitted to a level 1 trauma center sustained a PNI (Noble et al., 1998), highlighting the need for strategies that augment this inadequate regenerative response.

Phosphatase and tensin homolog (PTEN) is an inhibitor of the PI3K/AKT/mTOR cell growth and proliferation pathway (Stambolic et al., 1998). The loss of PTEN function *via* viral Cre transfection induces a regenerative response in corticospinal neurons after a spinal cord injury (K. Liu et al., 2010), a finding that has been reported in other CNS axonal injury models (Park et al., 2008). In the PNS, in which regeneration is more robust but still suboptimal, PTEN deletion has been shown to have modest augmentative effects on axonal regeneration (Gallaher and Steward, 2018). This allows for the unique opportunity to compare a pathology-dependent Cre upregulation strategy with a more traditional viral Cre transfection model.

Here, we present a novel transgenic mouse model that allows for the selective genetic editing of injured neurons via the insertion of a Cre-ERT2 construct under the native ATF3 promoter. We confirm expression of the Cre-ERT2 construct in injured neurons after PNI using a fluorescent reporter line and we demonstrate that this model may also be used to effect recombination in select populations of axotomized CNS neurons. Finally, we show that neuronal injury-specific Cre-ERT2 expression can be used to functionally alter the regenerative capacity of neurons though excision of crucial PTEN exons, a proof-of-principle experiment that illustrates the utility and potential of this model in neurotrauma.

Materials and Methods

Animals

All animal procedures were performed in accordance with the University of British Columbia and King's College London animal care committees' regulations. All mice were between two and four months of age and equally distributed between sexes.

The ATF3-CreERT2 (ATF3^{cre}) strain used was generated and described by Denk et al. (2015). Briefly, the Cre-ERT2 construct was inserted directly after the ATG start codon of the second ATF3 exon followed by a 3' untranslated region and a polyadenylation tagging sequence. For some experiments, this line was subsequently crossed to a floxed stop tdTomato Ai14 reporter line (RRID:IMSR_JAX:007908; Madisen et al., 2010). The ATF3^{cre} line was also crossed with conditional PTEN deletion line (PTEN^{fl/fl}) with LoxP sites flanking exon 5 of the



PTEN gene (RRID:IMSR_JAX:006440; Lesche et al., 2002; in some cases also crossed with the Ai14 line to determine recombination efficiency). Mice were maintained on a mixed C57BL/6J x 129SvEv background. ATF3-CreERT2 mice are available by request from the laboratories of Franziska Denk (King's College London) and Matt Ramer (the University of British Columbia).

Surgical procedures

For the sciatic nerve crushes the animals were administered buprenorphine (0.02 mg/kg; Temgesic) and ketoprofen (5 mg/kg; Anafen) subcutaneously for prophylactic analgesia. Once anaesthetized with isoflurane (5% induction, 2-3% maintenance; Fresenius Kabi Canada Ltd.), the sciatic nerve was exposed by blunt dissection and crushed for 15 s (thrice, 5 s each) with fine #5 forceps at the sciatic notch. For the brachial nerve crush the same anesthetic and analgesic protocols were used. The distal brachial plexus was exposed in the upper forelimb, the median radial, and ulnar nerves were crushed with fine forceps. Tamoxifen (Sigma) was dissolved in wheat germ oil (Denk et al., 2015) and injected at a concentration of 75 mg/kg at the time of injury. Pure anti-estrogen ICI 182780 (Tocris) was delivered by gavage the day before, the day of, and the day after injury (20 μ g dissolved in sunflower oil). For the 2-d regeneration assays, the injury site was marked with forceps dipped in graphite (ThermoFisher). For retrograde tracing, 1 μ l of 5% Fluorogold (Santa Cruz Biotechnologies) dissolved in 50:50 DMSO:PBS (Sigma-Aldrich) was injected intraneurally immediately before injury with a $10-\mu$ l Hamilton syringe (Sigma-Aldrich).

Spinal hemisection was also conducted with the same anesthetic and analgesic protocols outlined above. A midline incision was made over the lower cervical/upper thoracic spinal cord, and the C8 and T1 laminae were removed. A 25-gauge needle was inserted dorso-ventrally at the midline between the C8 and T1 spinal segments to allow relatively atraumatic entry of one blade of a pair of microscissors. The cord was laterally transected with microscissors; after hemostasis was established, the muscle and skin were closed in layers with sutures. Mice were killed 7 d later.

Functional outcomes

To investigate the following anatomic and functional recovery outcomes we employed a transgenic line with only the *Atf3* and *Pten* alleles altered (i.e., without reporter to avoid potential confounds associated with high tdtomato expression). Specifically, in the first set of experiments we compared ATF3^{+/+}, ATF3^{+/cre}, and ATF3^{cre/cre} mice to determine similarities to previous knock-out models (Gey et al., 2016). In the second, set we compared ATF3^{+/cre}:PTEN^{+/+} mice with ATF3^{+/cre}:PTEN^{fl/fl} to compare axotomy-induced PTEN deletion with virally-mediated PTEN deletion (Gallaher and Steward, 2018).

Behavioral testing

Reflex withdrawal or crossed extension (i.e., a nocifensive response) on strong toe pinch was assessed in mice lightly anesthetized with 2% isoflurane. On loss of righting reflexes, each animal was removed from the induction

chamber and placed prone on a table. To confirm light anesthesia, the base of a contralateral toe was pinched with curved serrated forceps. If there was no initial response, the other toes on the same foot were pinched in succession until either a response occurred, or there was an escape attempt. If a nocifensive response occurred, the ipsilateral toes were pinched starting with the first and ending with the fifth digit. As mice are prey species, they are prone to thanatosis (playing dead), and can suppress nocifensive withdrawal as they emerge from anesthesia. As such, a contralateral toe was pinched again following ipsilateral toe trials. An absence of a contralateral nocifensive response was almost always followed within 5 s by escape, and so the test was repeated. The test was also repeated if the mouse regained consciousness while the ipsilateral paw was being assessed. The presence or absence of a response to each ipsilateral toe pinch was recorded. If pinch to a particular toe elicited a response 2 d in a row, recovery was assigned to the first.

For the grasping assay the mouse was suspended upside-down from a wire cage lid and the grasping ability of the hindpaw was assessed and scored. Scores were assigned using the following semi-quantitative metric: undirected paw placement = 0, directed paw placement = 1, occasional grasp = 2, consistent grasp = 3. Both tests were conducted every other day starting on the third postoperative day until the experimental endpoint (day 28).

Electromyography

Four weeks following injury, mice were anesthetized with urethane (3 g/kg in dH₂O), and their sciatic nerves were exposed at mid-thigh. The nerves were bathed in paraffin oil and draped across a pair of silver wire hook electrodes (anode-cathode distance: 1 mm). EMG needle electrodes were placed subcutaneously over the lateral aspect of the hindpaw; one at the calcaneus, the other just proximal to the base of the 5th digit (over the abductor digiti minimi muscles). The nerve stimulated with 200 μs square wave current pulses using a stimulus generator (Master 9, A.M.P.I.) and stimulus isolator (A.M.P.I.). Signals were amplified using a Dual Bio-amp connected to a 16 channel Powerlab (ADInstruments). Signals were sampled at 40 kHz and filtered using LabChart7 software. After establishing appropriate electrode polarity, current pulses were delivered at increasing intensities until an EMG signal became apparent, and threshold current was recorded. The maximum EMG signal was then obtained, and the latency and amplitude of the first positive peaks were recorded.

Tissue processing

Mice were transcardially perfused with phosphate buffered saline followed by 4% paraformaldehyde (Thermo-Fisher). Once dissected, tissue was postfixed overnight in 4% paraformaldehyde, then overnight again in 20% sucrose (Thermo-Fisher) in 0.1 M phosphate buffer. The tissue was then frozen in Cryomatrix (Thermo-Fisher) and sectioned at 20 μm (DRG and sciatic nerve) or 50–100 μm (spinal cord). In some cases (Fig. 1), whole sympathetic or sensory ganglia were stained and imaged. All sections



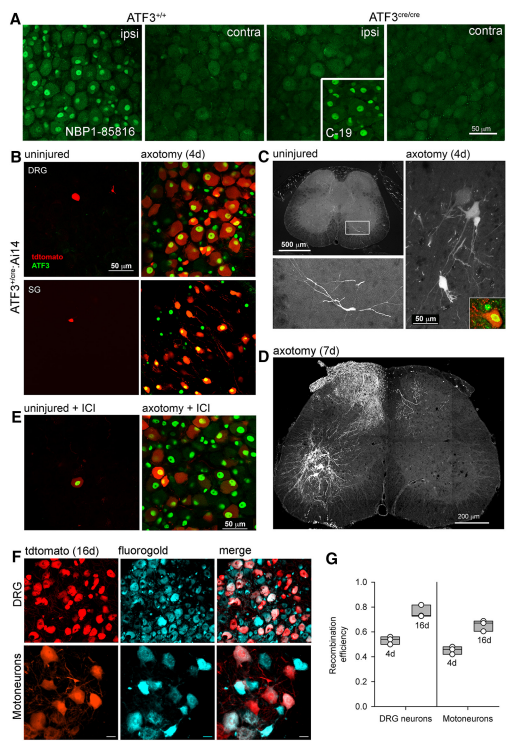


Figure 1. Axotomy-induced recombination in peripherally-projecting neurons. **A**, Validation of an ATF3-specific antibody. The Novus antibody (NBP1-85816) produces a positive signal in nuclei of axotomized sensory neurons in ATF3^{+/+} mice, but not ATF3^{cre/cre} mice. Note that the Santa Cruz antibody (C-19) labels neuronal nuclei in in the latter (inset), indicating non-specific staining. **B**, **C**, Axotomy induced reporter expression in sensory (DRG), sympathetic (stellate ganglion, SG), and motoneurons 4 d after injury. **D**, Reporter expression in sensory axons and motoneurons one week after injury. **E**, Preventing CreERT2 translocation from cytoplasm to nucleus with ICI 182780 reduces recombination in ATF3⁺ cells (by \sim 50%). **F**, Recombination efficiency 16 d after injury was calculated by expressing the proportion of tracer-filled somata (labeled at the time of injury) that were also reporter (tdtomato)-positive. **G**, Recombination efficiencies at 4 and 16 d after injury (n = 3 for each time point) for DRG and motoneurons. Images in panels **A**, **B**, **E** were taken from whole mounts, those in **C**, **D**, **F** from cryosections.



were blocked in 10% normal donkey serum with 0.2% Triton X-100 plus 0.02% sodium azide in PBS. Sections were incubated overnight with primary antibodies: rabbit anti-ATF3 (1:400, Santa Cruz SC-188), rabbit anti-ATF3 (1:500, Novus NBP 1-85816), rabbit anti-SCG10 (1:1000, Novus NBP 1-49461), rabbit anti-PTEN (1:400, Cell Signaling 9188), and mouse anti-PTEN (1:200, Cell Signaling 14642). Sections were incubated for 2 h with the appropriate secondary antibodies at a concentration of 1:1000: Alexa Fluor 488 donkey anti-rabbit (Invitrogen A21206), and Alexa Fluor 488 donkey anti-mouse (Jackson ImmunoResearch 715-545-151). Slides were cover-slipped with ProLong Gold with DAPI (Invitrogen).

Image acquisition and quantification

All images were acquired with a Zeiss LSM 800 confocal microscope using Zen (Blue) software. Recombination efficiency 4 d after lesion was determined by dividing the number of ATF3⁺-plus-tdtomato⁺ neuronal nuclei by the total number of ATF3⁺ neuronal nuclei. The 16-d recombination efficiency was calculated by dividing the number of tracer-positive neuron cell bodies by the total number of reporter-positive cell bodies. All image processing and quantification was done using ImageJ (Fiji version 2.0.0rc-66/1.52b).

Both PTEN antibodies produced specific staining, but with high background, varying depending on tissue examined (DRG and spinal cord). After comparing both antibodies across all tissues, our analyses relied on the Cell Signaling 9188 antibody for the DRG images and the Cell Signaling 14642 for the ventral root images. PTEN immunoreactivity in the DRG was quantified by measuring the mean pixel intensity of the entire cell layer of the DRG. In the spinal cord, PTEN immunoreactivity was weak in motoneurons, but intense in motor axons on either side of the ventral root exit zones, and so we focused our attention there. PTEN immunoreactivity in ventral root axons was determined by selecting for tdtomato⁺ axons and measuring the PTEN intensity of each axon.

To determine the regeneration density and distance along the sciatic nerve one best 20- μ m-thick section (i.e., lacking folds, bubbles or other sectioning artifacts) was selected for each animal, which was then fully imaged (z-stack and tiled). The stack was orthogonally projected into a single image through its entire depth. The images were then processed to generate binary overlays. The average density of SCG10 immunopositive axons along the width of the nerve was measured and then averaged over 100- μ m increments.

Statistics

All statistical analyses were conducted using Prism 7 (GraphPad). We used a one-way ANOVA followed by Tukey's multiple comparison test to compare PTEN immunoreactivity among DRGs (ipsilateral and contralateral from ATF3-CreERT2 mice with and without floxed PTEN. We used the Kolmogorov–Smirnov goodness-of-fit test to compare PTEN immunoreactivity in tdtomato+ axons between ATF3-CreERT2 tdtomato reporter mice with and without floxed PTEN. Differences in axonal regeneration between ATF3-CreERT2 mice with and without floxed

PTEN were determined on cumulative axon densities over 2-mm increments from the crush injury using either a one-way ANOVA followed by Dunnett's multiple comparison test (three groups) or an un-paired two-tailed t test (two groups). For toe pinch, we compared proportions of animals with five sensate digits (i.e., complete nocifensive recovery) over time following injury using a log-rank (Mantel–Cox) test. The same test was used to compare proportions of animals which showed consistent grasping with the injured hindpaw. For EMG data, we compared ipsilateral and contralateral values (threshold, latency, amplitude) within groups using paired t tests. Because we used males and females (which differ in size), for between-group comparisons we used unpaired t tests on ipsilateral/contralateral ratios.

Results

ATF3-driven injury-induced recombination

The Novus ATF3 antibody labeled the injured wild-type mice (ATF3^{+/+}) ipsilateral DRG neurons but did not label the same ipsilateral DRG neurons in the homozygous mutant mice (ATF3^{cre/cre}; Fig. 1A). However, the Santa Cruz antibody did label the injured ipsilateral DRG neurons in the ATF3^{cre/cre} mice, suggesting that it is not specific to ATF3 but another, injury-dependent protein as there was no neuronal labeling of any contralateral DRGs (Fig. 1A).

ATF3-driven recombination was exceedingly rare in the uninjured nervous system (Fig. 1*B*,*C*), although it was noted that uninjured recombination appeared to be more prevalent in older mice (data not shown). Four days after PNI, there was a robust tdtomato signal in the neuronal cell bodies in axotomized DRGs, stellate (sympathetic) ganglia (Fig. 1*B*), and ventral motor pools (Fig. 1*C*,*D*) in the ATF3^{+/cre} Ai14 reporter mice without the administration of tamoxifen. Tdtomato expression was only present in neurons, and only in those with nuclear ATF3 immunopositivity (Fig. 1*B*,*C*).

ATF3 expression is maximal at 4 d after injury but declines in sensory and motoneurons between 10 and 20 d (Tsujino et al., 2000). In 4-d lesions, we calculated the proportion of ATF3-positive DRG and motor neurons that were also tdtomato-positive (Fig. 1B,G). For 16-d lesions, we identified injured neurons by fluorogold labeling (injected at the time of injury), and determined the proportion that were also tdtomato labeled (Fig. 1*F*,*G*). In the DRG, recombination efficiencies were (mean \pm SEM) 53 \pm 2% (4 d) and 76 \pm 3% (16 d). For motoneurons, the efficiencies were 45 \pm 2% (4 d) and 65 \pm 3% (16 d). When the estrogen receptor α antagontist (ICI) was administered the amount of neuronal recombination was reduced by almost 50% (from 53 \pm 2% to 28 \pm 1%, p < 0.05, n = 3mice per group; Fig. 1E), demonstrating that recombination was due to leak of the CreERT2 construct into the nucleus as a result of inadequate cytoplasmic anchoring.

In the injured distal stump of the sciatic nerve, the Novus ATF3 antibody showed a uniform and punctate ATF3 signal that was attributable to white blood cells (Fig. 2A). When only the secondary antibody was used the same signal was present and not localized to the nucleus



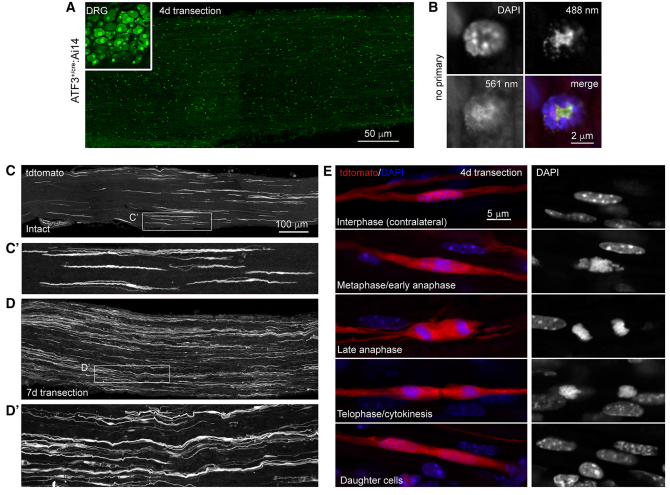


Figure 2. Axotomy does not induce ATF3 in Schwann cells. **A**, Cryosections from injured DRG (inset) and distal sciatic nerve from the same mouse processed for ATF3 immunohistochemistry (Novus NBP1-85816). **B**, Punctate staining in the nerve proved to be non-specific fluorescence of leukocytes (note non-nuclear signal in the absence of primary antibody). **C**, In intact sciatic nerves, cells morphologically identical to Remak cells had at some point undergone recombination. **D**, Following injury, their numbers increased. **E**, This was attributable to their proliferation in the injured nerve (as opposed to ATF3 induction and subsequent recombination). **C**', **D**', Magnification of areas outlined in **C** and **D** demonstrate the spindle shaped morphology characteristic of Remak cells.

of the putative white blood cells indicating that the antibody signal was an artifact, and that there was no ATF3 upregulation in the injured sciatic nerve (Fig. 2B).

In the uninjured sciatic nerve, long, spindle shaped cells that morphologically resemble Remak Schwann cells (RSCs; Gomez-Sanchez et al., 2017) were tdtomato+ and therefore had expressed ATF3 at some point in their lifetime (Fig. 2C). Seven days after axotomy, there was an increased density of these tdtomato+ RSCs (Fig. 2D), although they were not positive for ATF3 immunolabeling. On closer inspection 4 d after injury, we found multiple examples of tdtomato+ RSCs undergoing all phases of mitosis (Fig. 2E), suggesting that the increased density was not due to an injury induced upregulation of ATF3 but instead was the result of proliferation of previously-labeled cells.

ATF3's role in peripheral regeneration and functional recovery

Before employing the ATF3^{cre} line to edit the genes of injured neurons to manipulate the regenerative response

we first wanted to characterize the effect of losing one or both copies of the *Atf3* allele. Since the Cre insert prematurely terminates the ATF3 coding sequence, the ATF3^{cre} allele is notionally non-functional.

To determine whether loss of the *Atf3* allele attenuated functional recovery after PNI we examined behavioral recovery up to 28 d following sciatic crush, and electromyographical (EMG) activity at the experimental endpoint. We found that for both the pinching and grasping assays that the ATF3^{cre/cre} group had significantly diminished functional recovery (p = 0.0046 for pinch, p = 0.0189 for grasp; Fig. 3*A*,*B*). The performance of ATF^{cre/+} mice was intermediate for both assays (p = 0.0012 for pinch, p = 0.0049 for grasp) suggesting a gene dosage effect (the amount of functional transcript affects the level of recovery).

In terminal EMG experiments, we found that the ATF3^{cre/cre} group had significantly longer peak latencies (p = 0.0003; Fig. 3E), and smaller compound muscle action potentials (p = 0.002; Fig. 3F) when comparing ipsilateral/contralat-



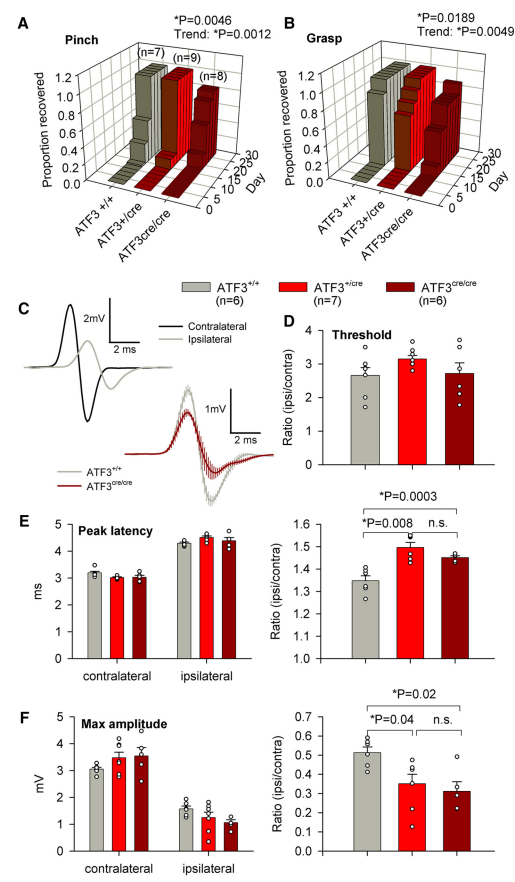


Figure 3. Loss of ATF3 function delays functional recovery following sciatic nerve crush. A, B, The rate of functional recovery



(nocifensive reflex withdrawal to a toe pinch and presence of any grasping ability) was reduced in mice lacking both wild-type ATF3 alleles (log rank Mantel–Cox test). Haploinsufficiency was also suggested by the statistically-significant trend from wild-type to homozygous knock-in (log rank Mantel–Cox test), n=7, n=9, and n=8 for ATF3^{+/+}, ATF3^{+/cre}, and ATF3^{cre/cre}, respectively. \bf{C} , Representative EMG traces from ipsilateral and contralateral sides of an ATF3^{+/+} mouse 28 d after sciatic nerve crush, and composite traces from seven ATF3^{+/+} mice and seven ATF3^{cre/cre} mice. \bf{D} , EMG thresholds did not differ between genotypes (paired t test). \bf{E} , \bf{F} , While absolute latencies and amplitudes did not differ between genotypes (paired t test), their ipsilateral/contralateral ratios (correcting for mouse size) indicated reduced conduction velocity (\bf{E}) and extent of reinnervation (\bf{F}) in mice lacking one or both wild-type ATF3 alleles (unpaired t test), n=6, n=7, and n=6 for ATF3^{+/+}, ATF3^{+/cre}, and ATF3^{cre/cre}, respectively.

eral ratios to the controls, indicating more complete muscle reinnervation in ATF3 $^{+/+}$ mice. There were no differences in activation thresholds (Fig. 3D). Surprisingly the ATF $^{\rm cre/+}$ group resembled the ATF $^{\rm cre/cre}$ group in EMG measures, displayed significantly longer peak latencies (p=0.008; Fig. 3E), and smaller compound muscle action potentials (p=0.04; Fig. 3F) when comparing ipsilateral/contralateral ratios to the controls providing further evidence of a gene dosage effect.

The above data unequivocally demonstrate delayed functional recovery (which can only be attributable to regeneration of injured axons given the absence of ATF3 expression in uninjured neurons) in ATF3-deficient mice. We then asked when differences in regeneration could be discerned between genotypes anatomically. To this end we assayed axonal regeneration along the sciatic nerve 2 and 3 d after crush by taking a single section (one that lacked sectioning artifacts like tears or folds) and imaged through its full 20-µm depth) for analysis. Two days after sciatic nerve crush, cumulative axon density was not yet different between ATF3+/+, ATF3cre/+, and ATF3cre/cre groups (Fig. 4A,B). Three days after sciatic nerve crush, a significant difference was detected in cumulative axonal density at 2-4 mm distal to the injury site between ATF3^{+/+} and ATF3^{cre/cre} groups (p = 0.0200), and the ATF3^{+/cre} group tested positive as a significant intermediary between groups (p = 0.0108; Fig. 4C,D). These anatomic results further support our findings that loss of one or both copies of the ATF3 allele diminishes regeneration after PNI.

ATF3-driven injury-induced PTEN knock-down

To determine whether the ATF3+/cre line can excise floxed endogenous genes, we crossed it to a floxed Pten line, crushed the sciatic nerve, and measured PTEN expression 28 d after injury. Three sections each from five animals were used for analysis. High background with both PTEN antibodies rendered precise estimates of recombination efficiency difficult, but the antibody labeled small-diameter DRG neurons, as reported previously by Gallaher and Steward (2018). There was no reduction in PTEN immunoreactivity between the ipsilateral and contralateral DRG cell layers in control animals (Fig. 5A) indicating that injury itself does not change PTEN expression. For reasons unknown, but possibly due to the leaky Cre-ERT2 construct, there was a significant difference in intensity measurements between genotypes on the contralateral (uninjured) side. Nevertheless, there was a significant reduction in PTEN expression on the ipsilateral side compared to the contralateral side of ATF3 $^{+/cre}$ PTEN $^{fl/fl}$ mice (p=0.0113) indicating that ATF3 driven Cre expression successfully excises the PTEN gene (Fig. 5A,B). The non-specific background PTEN antibody intensity was determined from sections from ATF3 $^{+/cre}$ PTEN $^{+/+}$ mice in which PTEN-positive DRG neuronal somata were excluded from regions of interest (i.e., the ROIs were the negative of the PTEN-positive neurons). PTEN immunoreactivity in ATF3 $^{+/cre}$ PTEN $^{fl/fl}$ mice was no different from background.

Spinal motoneurons were weakly PTEN-immuno-reactive, and although somata were less readily identifiable if they were also tdtomato-positive (Fig. 5C), background staining precluded reliable analysis in the ventral horn. We therefore examined PTEN expression in the ipsilateral ventral roots (which were intensely PTEN-immunoreactive on either side of the ventral root exit zone) 7 d after sciatic nerve crush in reporter mice with ATF3 $^{+/\text{cre}}$ and PTEN $^{\text{fl/fl}}$ or PTEN $^{+/+}$ alleles. Four control animals and three experimental animals were used (three sections from each). We found that PTEN expression was significantly reduced (p < 0.0001) in the axons that also expressed tdtomato in the PTEN $^{\text{fl/fl}}$ group compared to controls (Fig. 5D).

ATF3-driven PTEN excision improved functional recovery

To determine whether the ATF3^{+/cre} driven *Pten* excision was robust enough to effect enhanced functional recovery after PNI we examined behavioral recovery over a month following sciatic crush, and EMG activity at the experimental endpoint. For behavioral analysis the control group had 14 animals in it and the experimental group had nine. There were no differences in sensory (pinch) or sensorimotor (grasping) assays between the ATF3^{+/cre} PTEN^{fl/fl} and control groups (Fig. 6*A*,*B*), as has been reported previously (Gallaher and Steward, 2018).

In terminal EMG experiments, however, we found that the ATF3^{+/cre} PTEN^{fl/fl} group had a significantly lower EMG activation threshold (p < 0.0001; Fig. 6*E*), shorter peak latencies (p = 0.040; Fig. 6*F*), and larger compound muscle action potentials (p = 0.003; Fig. 6*G*) when comparing ipsilateral/contralateral ratios to the controls, indicating more complete muscle reinnervation in mice lacking PTEN.

Enhanced functional recovery in PTEN deficient mice implies more robust axonal regeneration. We were again curious as to when axonal regeneration following ATF3-driven injury induced *Pten* excision might be detected histologically following sciatic crush injury. Six animals



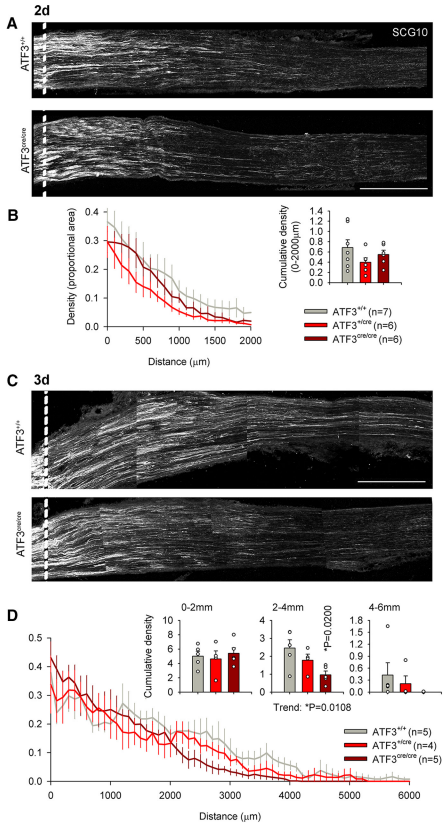


Figure 4. Loss of ATF3 function modestly reduces axonal regeneration following sciatic nerve crush. **A**, **B**, There was no difference in axonal regeneration 2 d following injury between ATF3^{+/+}, ATF3^{+/cre}, and ATF3^{cre/cre} mice (n = 7, n = 6, and n = 6, respectively, groups were compared with a one-way ANOVA on cumulative densities).**C**,**D**, ATF3^{cre/cre} mice exhibited significantly diminished axonal regeneration 3 d following injury 2–4 mm distal to the injury compared to ATF3^{+/+} mice <math>(n = 5 for both groups, one-way)



ANOVA followed by Dunnett's multiple comparison test). The hemizygous group (n=4) tested positive as a significant intermediary between both control and ATF3 null groups (post hoc test for trend). Dotted line indicates distal border of crush site, 500 μ m from the edge of the block. Scale bars: 500 μ m.

were used in each group and a single section (lacking sectioning artifacts, and imaged through its full 20- μ m depth) was taken for analysis. Two days after sciatic nerve

crush, axon density plotted as a function of distance was obviously increased in ATF3 $^{+/\rm cre}$ PTEN $^{\rm fl/fl}$ over that in ATF3 $^{+/\rm cre}$ PTEN $^{+/+}$ mice, and the cumulative density at 2

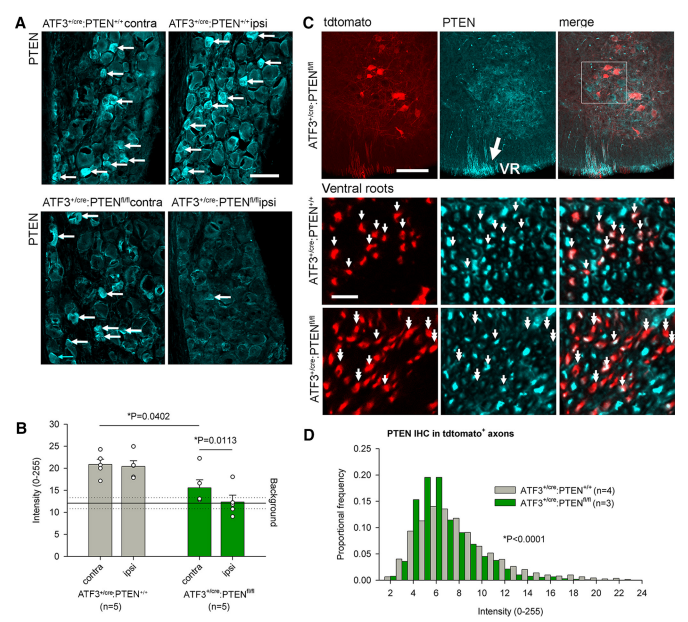


Figure 5. Axotomy-induced PTEN deletion in sensory and motoneurons. **A**, **B**, Axotomy results in significant loss of PTEN expression in the DRG. Arrows in **A** indicate small PTEN-positive DRG neurons. The PTEN antibody results in high background staining in all animals, to which PTEN immunoreactivity is reduced in axotomized DRGs of ATF3^{+/cre}:PTEN^{fl/fl} mice (**B**; one-way ANOVA followed by Dunnett's multiple comparison test, n=5 for both groups). **C**, PTEN immunoreactivity is weak in all motoneurons, rendering difficult confirmation of axotomy-induced knock-down. However, ventral root (VR; large arrow) axons close to the ventral root exit zone were intensely immunopositive, single arrows indicate axons that were both tdtomato and PTEN positive, whereas double arrows indicate recombination without PTEN immunoreactivity. In sections of ventral roots, we were able to demonstrate a significant decrease in PTEN immunoreactivity in tdtomato-positive axons (**D**; Kolmogorov–Smirnov goodness of fit test), n=4 and n=3 for ATF3^{+/cre}:PTEN^{+/+} and ATF3^{+/cre}:PTEN^{fl/fl}, respectively. Scale bars: 50 μm (**A**), 100 μm (**C**), 10 μm (**E**).



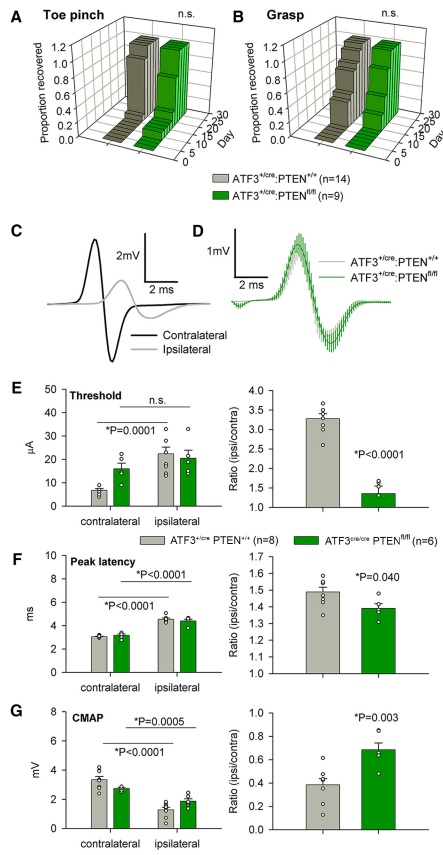


Figure 6. Axotomy-induced PTEN deletion and improves functional recovery following sciatic nerve crush. \bf{A} , \bf{B} , While there was no difference in recovery of reflex nociception or grasping in mice with floxed PTEN alleles (log rank Mantel–Cox test, n=14 and n=14) and n=14.



9 for ATF3 $^+$ /cre:PTEN $^+$ /+ and ATF3 $^+$ /cre:PTEN $^{fl/fl}$, respectively), EMG responses (**C–G**) indicated enhanced recovery of neuromuscular function. **C**, **D**, Representative EMG traces from ipsilateral and contralateral sides of an ATF3 $^+$ /cre:PTEN $^+$ /+ mouse 28 d after sciatic nerve crush (**C**) and composite traces from eight ATF3 $^+$ /cre:PTEN $^+$ /+ mice and six ATF3 $^+$ cre:PTEN f /fl mice. **E–G**, Ipsilateral/contralateral ratios of EMG thresholds (**E**), peak EMG latencies (**F**), and maximum CMAP amplitudes (**G**) all indicated more complete muscle reinnervation 28 d after injury; n=8 and n=6 for ATF3 $^+$ /cre:PTEN $^+$ /+ and ATF3 $^+$ /cre:PTEN f /fl, respectively, averages were compared using paired t tests and ipsilateral/contralateral ratios with unpaired t tests.

mm was statistically greater (p = 0.0008; Fig. 7). This demonstrates that the ATF3^{+/cre} line is able to edit the genes of injured neurons with enough efficacy to produce anatomic differences in regeneration.

ATF3-driven recombination after CNS injury

After confirming that the ATF3^{cre} line is capable of efficient recombination that can selectively edit genes the genes of injured neurons after PNI we wanted to determine whether the same line was applicable to CNS injury. Six ATF3^{cre/+}:Ai14 reporter mice underwent a C8/T1 lateral spinal hemisection, were killed 7 d after injury, and recombination throughout the entire CNS was characterized. In the spinal cord, several tracts were reliably labeled, these included putative rubrospinal, raphespinal, reticulospinal, and vestibulospinal descending tracts (Fig. 8C,D). Additionally, ascending neurons were tdtomato+; based on anatomic position these are likely to be spinothalamic spinocerebellar neurons (Fig. 8E,F). In the cervical cord, rostral to the injury site, there was a substantial amount of recombination in primary afferents innervating the dorsal horn (Fig. 8C). Four deep brain nuclei were consistently reporter-positive, these were the rubrospinal, reticulospinal, vestibulospinal, and paraventricular hypothalamic nuclei (Fig. 9A–E). There were examples of ATF3 immunolabeled and tdtomato-positive neurons in each of these nuclei (Fig. 9F).

Discussion

The expression of Cre recombinase has for the most part been restricted to either developmentally distinct subpopulations of cells via its insertion under specific promoters or to distinct physical regions by viral transfection. Selective cellular genetic modification of neurons based on functional state is an attractive and important refinement to this approach. Guenthner et al. (2013), for example, used a genetic labeling technique controlled by neuronal activity. In this case, recombination was driven by promoters for the immediate-early genes Fos and Arc, upregulated as part of the "excitation-transcription" neuronal response to synaptic activity initiated by CREB phosphorylation. Here, we present a novel transgenic mouse line that expresses Cre only once peripherallyprojecting neurons have been axotomized by inserting its construct into the native ATF3 locus. We demonstrate a substantial amount of recombination (~50% by 4 d after

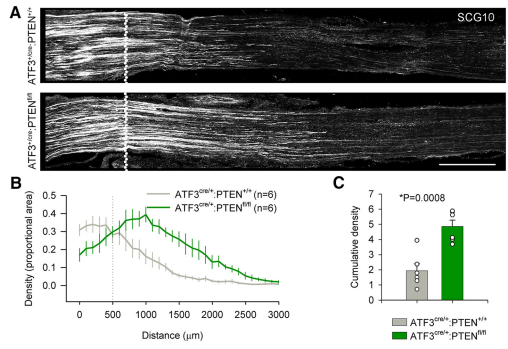


Figure 7. Axotomy-induced PTEN deletion and anatomic regeneration following sciatic nerve crush. **A**, **B**, Axonal regeneration 2 d subsequent to sciatic nerve crush (dotted line) was significantly enhanced in ATF3^{+/cre}:PTEN^{fl/fl} mice. In **C**, bar graphs represent the cumulative density of SCG10 immunoreactivity from 0 μ m (the distal extent of the crush site) to 2000 μ m; n=6 for both groups and cumulative densities were compared using an unpaired t test. Scale bars: 500 μ m.



A ATF3+/cre:Ai14

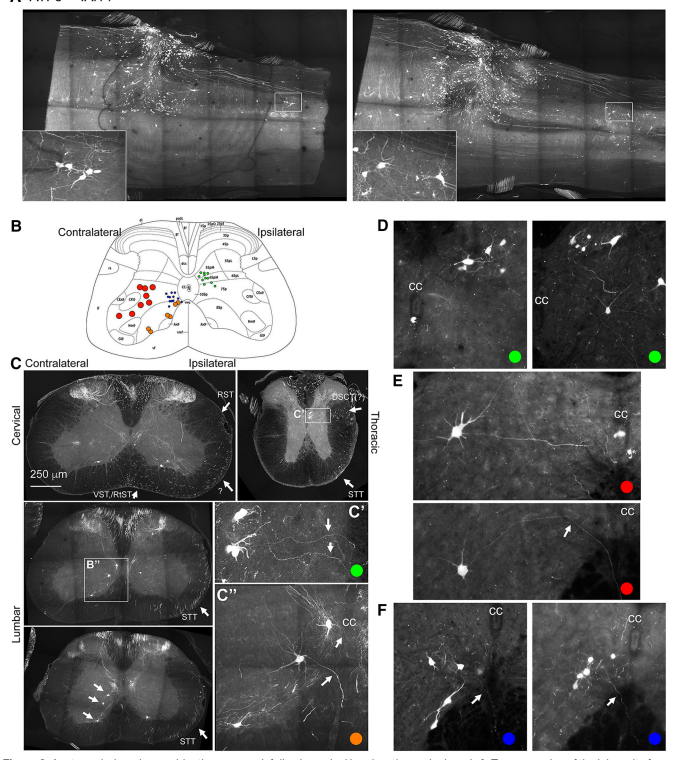


Figure 8. Axotomy-induced recombination one week following spinal hemisection: spinal cord. **A**, Two examples of the injury site from separate animals in longitudinal section. Insets show consistently-recombining ipsilaterally-projecting neurons near Clarke's column below the injury. **B**, Cartoon of transverse section of the hemisected spinal cord illustrating relative positions of positionally and morphologically distinguishable recombined neurons (colored dots correspond to examples in **C-F**). **C**, Examples of recombination after injury in cervical (top left), lumbar (two examples middle and bottom-left), and thoracic (top right). The most consistent findings were small ipsilaterally-projecting neurons in the thoracic cord (**C'**), and large neurons contralateral to injury from just lateral to area X to the ventral gray matter in the lumbar cord (**C''**). Arrows in **C'**, **C''** indicate midline-crossing axons. Arrow pointing to tdtomato⁺ axons in dorsal cervical white matter indicates probable rubrospinal (RST) and/or raphespinal tracts. Arrows pointing to reporter-positive axons in ventral white matter indicate



probable reticulospinal (RtST) and vestibulospinal (VST), and an unknown descending projection (?) tracts rostral to the injury, and spinothalamic (STT) and possible dorsal spinocerebellar [DSCT(?)] below the injury (thoracic and lumbar sections). **D**, Neurons in Clarke's column ipsilateral to the hemisection. **E**, **F**, Large and small, respectively, putative spinothalamic tract neurons contralateral to the hemisection. Arrows in **E**, **F** indicate commissural axons; cc, central canal.

injury, rising to \sim 65–80% by 16 d) in injured sensory, motor, and sympathetic neurons that is selective to axotomy. This injury-dependent Cre expression is restricted to the neurons

that have been axotomized, is rare in uninjured controls, and absent in peripheral glial cells. Furthermore, we show that axotomy-induced Cre expression can excise floxed genes

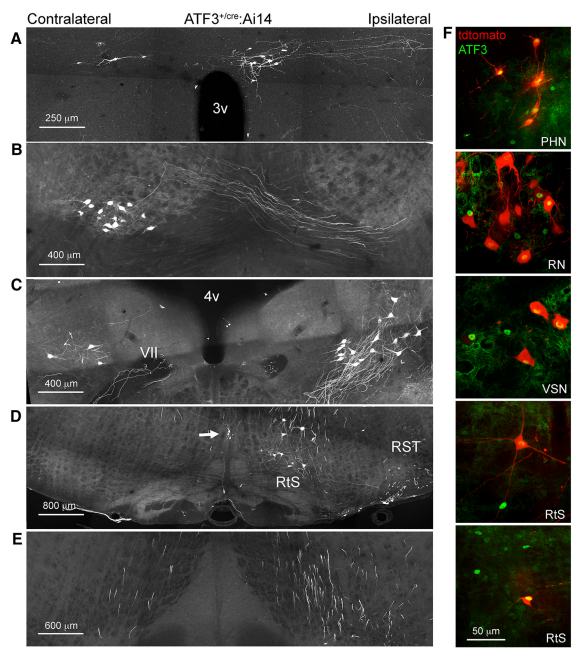


Figure 9. Hemisection-induced recombination in supraspinal neurons one week after injury. **A**, Paraventricular hypothalamic nucleus, descending part. **B**, Red nucleus. **C**, Vestibulospinal nucleus (the genu of the facial nerve is indicated by VII). **D**, Reticulospinal neurons (RtS) and raphespinal neurons (arrow), and the rubrospinal tract (RST). **E**, Reporter-expressing axons in the medial longitudinal fasciculus (conveying descending projections of reticulospinal and vestibulospinal axons). **F**, Examples of ATF3-positive, tdtomatopositive and tdtomato-negative neurons in the paraventricular hypothalamic nucleus (PHN), red nucleus (RN), vestibulospinal nucleus (VSN), and reticulospinal neurons (RtS).



with sufficient efficacy to significantly effect anatomic regeneration and functional recovery.

It is important to note that despite the Cre recombinase being anchored to a mutated estrogen receptor (ERT2) we have achieved significant recombination without the administration of tamoxifen. This can be attributed to the "leakiness" of the ERT2 construct where the ERT2 protein overwhelms its cytoplasmic anchor and translocates to the nucleus without tamoxifen (or its metabolites) binding. The high degree of tamoxifen-independent recombination we report is likely driven by massive upregulation of ATF3 in the PNS after injury: the *Atf3* gene (and hence Cre-ERT2) is highly transcribed once the neuron is axotomized, and there is a greater likelihood of ERT2 leak and subsequent recombination. Reduction of recombination by ICI 182780 (Fig. 1) provides an opportunity to titrate recombination due to leakiness.

We did not use tamoxifen to improve the recombination efficiency of the model, as it has been previously reported that tamoxifen administration, at the standard dose of 75 mg/kg, upregulates ATF3 without injury (Denk et al., 2015). Notionally it is possible that a lower dose of tamoxifen would not induce ATF3 expression, but would still improve the recombination efficiency after injury, although this dose has yet to be determined. Regardless, the efficiencies achieved in this report due to ERT2 leak remain sufficient to excise *Pten* and significantly improve anatomic regeneration and functional recovery.

The potential utility of our model is strengthened by similarities to previous manipulations using different techniques. A study by Gallaher and Steward (2018) investigated the effect of Pten deletion following axotomy in the sensory neurons that innervate the sciatic nerve. They excised Pten through a more traditional intraganglionic injection of an AAV Cre vector into the L4 and L5 DRG. Their study and ours agree on three key results: (1) PTEN immunohistochemistry preferentially labels small diameter sensory neurons, (2) PTEN deletion increased axonal regeneration along the sciatic nerve 3 d after axotomy, and (3) PTEN deletion after sciatic nerve crush did not significantly improve sensory functional recovery. The ATF3-CreERT2 line has several distinct advantages over more traditional Cre delivery models; the technical viral transfection setup is not necessary and Cre can be expressed in neuronal populations that may not be amenable to local injection. Moreover, the ATF3-ERT2 line allows for the selective editing of only injured neurons where a viral Cre transfection does not preclude the possibility of Cre expression in uninjured neurons.

Using a homozygous knock-in (i.e., null mutant), we tested the validity of two reported ATF3-specific polyclonal antibodies: Novus (NBP 1-85816) and Santa Cruz (C-19). ATF3 signal was expectedly absent in the ATF3 null ipsilateral DRG after axotomy when using the Novus antibody. However, signal was still present in both the control and ATF3 null ipsilateral (but not contralateral) DRG when tested with the Santa Cruz antibody suggesting that is not specific to ATF3, but likely another injury-dependent protein. This could call into question some of the conclusions made by the over 150 articles that have

used this antibody. We then used the Novus ATF3 antibody to investigate ATF3 expression in the injured sciatic nerve. Despite the appearance of putative ATF3 upregulation, we were able to determine that this signal was restricted to white blood cells, non-nuclear, and in fact due to background fluorescence. Clements et al. (2017) reports Schwann cell ATF3 mRNA expression in the uninjured sciatic nerve which is then reduced on injury, which appears to contradict our observations of a lack of ATF3-driven recombination in normal Schwann cells but supports our finding of a lack of injury-dependent glial cell ATF3 expression. To obtain these data the Schwann cells needed to be dissociated, purified, and then sorted before sequencing. Given ATF3's role as a stress response immediate early gene it is certainly plausible that this baseline ATF3 expression is due to the FACS process and not in fact expressed in the *in vitro* uninjured nerve, this has indeed proven to be an issue in other cell types, such as muscle (van den Brink et al., 2017). Histologic evidence of ATF3 mRNA upregulation following nerve injury is also unconvincing (Hunt et al., 2004). We therefore conclude that ATF3 is not upregulated in peripheral glia after injury cells, contrary to previous reports. This finding further strengthens the usefulness of the model as any axotomydependent Cre upregulation is restricted to neuronal populations.

Tdtomato-positive RSCs increased in density following sciatic nerve injury. RSCs are a subtype of Schwann cell that are non-myelinating but still ensheathe small caliber axons to form Remak bundles (Harty and Monk, 2017). Gomez-Sanchez et al. (2017) characterized this subtype of Schwann cells using sporadic permanent fluorophore labeling. These cells are spindle shaped, can be branched, and are $\sim\!250~\mu\mathrm{m}$ in length; all of which are characteristics of the sciatic nerve cells found to undergo injury-independent recombination in the ATF3^{cre} Ai14 reporter line. The absence of ATF3 immunoreactivity in these cells, along with the high numbers of RSCs that can be observed undergoing mitosis 4 d after injury show that their increase in density is due to cellular division rather than injury-induced ATF3-cremediated recombination.

The ATF3^{cre/cre} null mutant also allowed us to test the effect of a loss of ATF3 function in PNS regeneration early after injury. While we found no significant differences between the groups in axonal regeneration along the sciatic nerve at the 2-d time point, we did find a statistically significant functional deficit of both ATF3-deficient groups in both our behavioral and EMG assays. Despite being a standard time point for this type of analysis, the lack of significantly different 2-d axonal regeneration along the sciatic nerve could be because the axons have not had sufficient time to differentially regenerate enough to produce a measurable difference. This supports previous work done by Gey et al. (2016), who found attenuated regeneration after facial nerve axotomy in ATF3 null mice. Gey et al. (2016) also showed that when DRG neurons were cultured in the presence of NGF, any differences in outgrowth between wild-type and knock-out neurons ATF3 were abolished. NGF is produced early on in the distal transected nerve (Heumann et al., 1987), and this early abundance may com-



pensate for any ATF3-mediated regenerative differences. By 3 d after injury, however, a clear difference between ATF3^{+/+} and ATF3^{cre/cre} mice had emerged, along with evidence again for haploinsufficiency (the intermediate position of ATF3^{+/cre} mice). To our knowledge, this is the first evidence that a loss of ATF3 mitigates functional recovery after injury.

Interestingly the existing literature suggests that ATF3 heterozygotes are haplosufficient (Gey et al., 2016) which is contrary to our findings of a gene dosage effect. The fact that the amount of ATF3 transcript affects the functional outcome after peripheral axotomy might seem surprising given how drastically ATF3 is upregulated in neurons after injury. However, given that ATF3 is a bZIP transcription factor that dimerizes with itself or other bZIP transcription factors to affect transcription, it is conceptually possible that the binding affinity for the ATF3 dimerization pair responsible for improving regeneration is low and therefore many copies of ATF3 must be produced for its pro-regenerative effect. Recent work done be Rodríguez-Martínez et al. (2017) exemplified this by showing that ATF3's DNA binding site preference to be highly dependent on its bZIP dimerization.

ATF3 gene dosage effects (i.e., partial function in hemizygous mice) should not be an impediment to investigations of functions of other (floxed) genes or sequences provided the appropriate controls are used. There is simply a new baseline on which other genetic manipulations (using ATF3-Cre-driven excision of floxed gene "X") can be evaluated. For this reason it is important that any investigation of the function of gene X compares ATF3 $^{+/\text{cre}}$: X $^{+/+}$ with ATF3 $^{+/\text{cre}}$:X $^{fl/fl}$ (or ATF3 $^{+/\text{cre}}$:X $^{+/fl}$, for investigating possible haploinsufficiency of X), as we have done here substituting "X" with "PTEN."

ATF3 is not only upregulated after PNI but also after CNS trauma including spinal cord injury (Huang et al., 2007), albeit usually meagerly and/or transiently, rendering expression patterns difficult to reveal (for review, see Hunt et al., 2012). As such we wanted to determine whether CNS neurons could be accessed genetically via ATF3-cre-mediated recombination after a lateral spinal cord hemisection. We observed consistent recombination in four deep brain nuclei (rubrospinal, reticulospinal, vestibulospinal, and hypothalamic) and their projecting axons and in at least two putative ascending tracts (spinocerebellar and spinothalamic). There was no recombination observed in the injured corticospinal tract, but this is not surprising given that it takes intracortical axotomy to induce ATF3 in corticospinal neurons (Mason et al., 2003): the magnitude of upregulation of regeneration associated genes (of which ATF3 is) have been documented to be dependent on the distance from the axon transection to the soma (Fernandes et al., 1999).

The mammalian PNS is able to regenerate after injury, unlike the CNS where regeneration does not occur. This positions the PNS as an excellent model for study to better understand what is necessary for mammalian neuronal regeneration. The ability to edit the genes of injured neurons and further dissect what is necessary and/or sufficient to produce this regeneration is of obvious value, and is now possible with the ATF3^{cre} transgenic line. ATF3

is also expressed in injured neurons after spinal cord injury (Huang et al., 2007; Wang et al., 2015; Darlot et al., 2017), traumatic brain injury (Förstner et al., 2018), and ischemic stroke (Song et al., 2011), making the line useful to scientists interested in the neuronal response to each of those pathologies.

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