Removal of the Potassium Chloride Co-Transporter from the Somatodendritic Membrane of Axotomized Motoneurons Is Independent of BDNF/TrkB Signaling But Is Controlled by Neuromuscular Innervation

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Abstract
The potassium-chloride cotransporter (KCC2) maintains the low intracellular chloride found in mature central neurons and controls the strength and direction of GABA/glycine synapses. We found that following axotomy as a consequence of peripheral nerve injuries (PNIs), KCC2 protein is lost throughout the somatodendritic membrane of axotomized spinal cord motoneurons after downregulation of kcc2 mRNA expression. This large loss likely depolarizes the reversal potential of GABA/glycine synapses, resulting in GABAergic-driven spontaneous activity in spinal motoneurons similar to previous reports in brainstem motoneurons. We hypothesized that the mechanism inducing KCC2 downregulation in spinal motoneurons following peripheral axotomy might be mediated by microglia or motoneuron release of BDNF and TrkB activation as has been reported on spinal cord dorsal horn neurons after nerve injury, motoneurons after spinal cord injury (SCI), and in many other central neurons throughout development or a variety of pathologies. To test this hypothesis, we used genetic approaches to interfere with microglia activation or delete bdnf from specifically microglia or motoneurons, as well as pharmacology (ANA-12) and pharmacogenetics (F616A mice) to block TrkB activation. We show that KCC2 dysregulation in axotomized motoneurons is independent of microglia, BDNF, and TrkB. KCC2 is instead dependent on neuromuscular innervation; KCC2 levels are restored only when motoneurons reinnervate muscle. Thus, downregulation of KCC2 occurs specifically while injured motoneurons are regenerating and might be controlled by target-derived signals. GABAergic and glycineergic synapses might therefore depolarize motoneurons disconnected from their targets and contribute to augment motoneuron activity known to promote motor axon regeneration.

Key words: excitability; GABA; KCC2; nerve injury; neuromuscular junction; regeneration

Significance Statement
The neuronal potassium-chloride cotransporter KCC2 is dysregulated after numerous types of neuronal injuries and has been related to neuronal dysfunction, hyperalgesia and spasticity. In this study, we investigated KCC2 regulation on spinal motoneurons with axons injured during peripheral nerve transections. We illustrate that KCC2 loss on axotomized motoneurons occurs at the transcriptional level. This loss differs in time course, completeness and signaling mechanisms from KCC2 dysregulation in other neurons after various pathologies or in motoneurons after spinal cord injury (SCI). In contrast, KCC2 loss on axotomized motoneurons relates to muscle innervation, suggesting a dependence on target-derived signals. We argue that KCC2 loss in axotomized motoneurons may be part of the response that facilitates regeneration of motor axons in the periphery.
Introduction

Peripheral nerve injury (PNI) affects over 200,000 people annually in the United States and recovery outcomes are poor, including loss of sensorimotor function that can persist for years (Mundie, 1920; Taylor et al., 2008; Höke and Brushart, 2010). A major problem hindering recovery is the slow speed of regeneration of axons in the periphery (1–3 mm/d; Sunderland, 1947; Brashurt, 2011). Treatments that accelerate motor axon growth include exercise and electrical stimulation, both dependent on activity and the release of BDNF (Al-Majed et al., 2000a, b; English et al., 2007, 2009; Udina et al., 2011). Counterintuitively, there is also a loss of excitatory synapses on the cell bodies of axotomized neurons while inhibitory synapses are largely retained (Lindé et al., 1992, 2000; Brännström and Kellerth, 1998; Novikov et al., 2000; Alvarez et al., 2011; Rotterman et al., 2019). This change in excitatory/inhibitory balance is usually interpreted as a mechanism to counter hyperexcitability such that motoneurons enter a “quiet” phase, focusing resources in gene and protein expression to promote regeneration (Choi, 1992; Oliveira et al., 2004). Indeed, enhanced preservation of inhibitory synapses in axotomized motoneurons increases the speed of muscle reinnervation and recovery of motor function (Berg et al., 2012).

However, despite the common argument that inhibitory synapse preservation helps keep injured motoneurons quiescent, GABAergic and glycineric synapses on axotomized motoneurons may not provide effective inhibition. Isoform two of the potassium chloride cotransporter (KCC2) regulates intracellular chloride in most adult neurons; it defines the strength of inhibition (Payne et al., 2003; Kaila et al., 2014; Kahle and Delpire, 2016) and even the direction of membrane polarizations after opening of GABA_A and glycine receptor chloride channels. KCC2 expression is downregulated in brainstem motoneurons following peripheral axotomy (Nabekura et al., 2002; Toyoda et al., 2003; Tatetsu et al., 2012; Kim et al., 2018) and results in spontaneous membrane depolarizations dependent on GABA_A and NMDA receptor activation that trigger calcium entry through the opening of voltage-gated calcium channels (Toyoda et al., 2003).

KCC2 expression in spinal motoneurons after peripheral axotomy has not been investigated. Most studies on adult spinal motoneuron KCC2 relate to dysregulation after spinal cord injury (SCI), which results in disinhibition and behavioral spasticity (Boulenguez et al., 2010; Murray et al., 2011; Viemari et al., 2011; Bos et al., 2013). Studies investigating spinal KCC2 after PNI, however, have focused on KCC2 dysregulation in dorsal horn neurons associated with mechanisms that promote hyperalgesia (also by disinhibition; Coull et al., 2003, 2005; Ferrini and De Koninck, 2013). In all of these conditions, decreases in KCC2 are considered maladaptive. As such, pharmacological enhancement of KCC2 activity has been a focus for targeted treatments of spasticity and neuropathic pain (Gagnon et al., 2013; Kahle et al., 2014; Sánchez-Brualla et al., 2018). KCC2 loss after PNI or SCI was found to be rapid but incomplete, and was the consequence of altered KCC2 trafficking and/or stability at the membrane (Rivera et al., 2002; Bos et al., 2013; Kahle et al., 2013; Medina et al., 2014). In particular, a microglia–BDNF–TrkB pathway is often highlighted as responsible for downregulation of KCC2 in spinal neurons after PNI (Coull et al., 2005; Ferrini and De Koninck, 2013). Other studies also point to BDNF–TrkB signaling as responsible for regulating KCC2 levels during development and under a variety of disease conditions, acting on both membrane trafficking and gene expression (Lee-Hotta et al., 2019). In contrast to the above studies, spinal motoneurons are directly injured (axotomized) after PNI, and whether KCC2 is also lost and whether it involves similar mechanisms of BDNF and TrkB signaling is unknown. The mechanisms that downregulate KCC2 in brainstem motoneurons after PNI have not been elucidated.

In this study, we show that KCC2 is decreased in spinal motoneurons after PNI. This loss is more extensive than after SCI and occurs throughout the somatodendritic membrane, but proceeds with a relatively slow time course (weeks instead of hours) and after downregulation of KCC2 mRNA. Using genetic approaches and pharmacological manipulations, we also show that the mechanism is independent of microglia activation and BDNF or TrkB signaling. However, we observed an intriguing dependence of KCC2 recovery on neuromuscular junction (NMJ) innervation, indicating that target-dependent factors might be involved in KCC2 regulation. The tight link between KCC2 and regeneration state raises the possibility that the absence of KCC2 from disconnected motoneurons is an adaptive phenomenon related to regeneration mechanisms, an idea we further explore in the discussion.

Materials and Methods

Animal care, procedures, and euthanasia were performed under prior approval by the Institutional Animal Care and Use Committee of Emory University and in accordance with the NIH Guide for the Care and Use of Laboratory Animals.
Table 1. Mice used to derive experimental genotypes

| Mouse strain               | Stock no.          | References      |
|----------------------------|--------------------|-----------------|
| CX3CR1<sup>EGFP</sup>     | RRID:IMSR_JAX:005582 | Jung et al. (2000) |
| CX3CR1<sup>creER</sup>+    | RRID:IMSR_JAX:020940 | Yona et al. (2013) |
| CSF1<sup>ff</sup>          | Donated Dr. Jean X. Jiang, University of Texas, San Antonio, TX | Harris et al. (2012) |
| SLICK-A                    | RRID:IMSR_JAX:007606 | Young et al. (2008) |
| F616A                      | RRID:IMSR_JAX:022363 | Chen et al. (2005) |
| ChAT<sup>Thy1</sup>-res<sup>-cre</sup>+ | RRID:IMSR_JAX:006410 | Rossi et al. (2011) |

Animals

Several lines of transgenic animals in addition to wild types (WTs) of both sexes were used. Except where noted, mice were obtained from The Jackson Laboratory and maintained on a mixed C57Blk/6J and 6N background. Table 1 lists all transgenic mouse lines used, their JAX codes, and RRID numbers.

To label microglia, we used Cx3cr1<sup>GFP</sup>+ heterozygous mice that carry green fluorescent protein (GFP) replacing a single copy of the endogenous fractalkine receptor gene, thereby enabling visualization of microglia (Jung et al., 2000). CX3CR1 is expressed exclusively in microglia in the CNS and subsets of myeloid cells in the periphery (Mizutani et al., 2012). In Cx3cr1<sup>GFP</sup>GFP homozygous mice, both alleles are replaced; these animals lack CX3CR1 expression and display altered microglia function in several diseases (Limatola and Ransohoff, 2014).

To target microglia for cell-specific deletions of BDNF, we used tamoxifen-inducible cre, crossing Cx3cr1<sup>creER</sup>+ mice with mice carrying floxed <i>bdnf</i> alleles (<i>bdnf<sup>f/f</sup></i>). Although CX3CR1 is expressed in peripheral myeloid cells as well as microglia, peripheral myeloid cells have a higher turnover rate (Bennett et al., 2016). Thus, four weeks after tamoxifen treatment only microglia remain <i>bdnf</i> genetic knock-outs (KO), while CX3CR1-expressing myeloid cells have been newly generated from myeloid precursors that lack CX3CR1 expression and have therefore not undergone tamoxifen-induced cre recombination (Goldmann et al., 2013; Tay et al., 2017).

To prevent the microglia reaction after PNI, specifically in the ventral horn of the spinal cord, mice were studied in which the <i>csf1</i> gene for colony stimulating factor 1 (CSF1) was knocked out in motoneurons. CSF1 is typically released from injured neurons to activate and recruit microglia (Elmore et al., 2014; Guan et al., 2016). We crossed Chat<sup>REScre</sup>+ with <i>csf1<sup>ff</sup></i> mice to eliminate CSF1 release from cholinergic neurons, including axotomized motoneurons. This manipulation has been shown to greatly attenuate the ventral horn microglial response to PNI (Rotterman et al., 2019). Mice carrying <i>csf1<sup>ff</sup></i> alleles were generously donated by Dr. Jean X. Jiang (University of Texas, San Antonio, TX).

We also crossed Chat<sup>REScre</sup>+ animals with <i>bdnf<sup>f/f</sup></i> mice to remove BDNF expression from motoneurons after injury. In this case, the <i>bdnf</i> gene was deleted from motoneurons throughout development. To delete <i>bdnf</i> more specifically in adult motoneurons, we used tamoxifen-inducible single-neuron labeling with inducible CreER-mediated KO (SLICK) mice. Specifically, mice of the SLICK-A line were crossed to <i>bdnf<sup>f/f</sup></i> mice. SLICK-A mice express YFP and tamoxifen-inducible cre in subsets of neurons controlled by the <i>thy1</i> promoter (Young et al., 2008). When treated with tamoxifen, cre recombinase is activated in YFP+ neurons and eliminates expression of floxed genes. Thus, after tamoxifen treatment, YFP+ axotomized motoneurons (expressing cre) can be compared with YFP− axotomized motoneurons (not expressing cre) within the same animal (Young et al., 2008; Wilhelm et al., 2012; Zhu et al., 2016).

To investigate the difference between α and γ motoneurons, one animal underwent unilateral sciatic nerve cut and ligation without prior retrograde injections. γ Motoneurons are not typically labeled by retrograde tracers and general markers for motoneurons (ChAT) and neuronal injury (activating transcription factor 3, ATF3) in addition to cell size were used to identify and quantify KCC2 on different populations of injured motoneurons as described below.

Tamoxifen treatment

When using tamoxifen-inducible cre mice, the drug (0.75 mg/20 g body weight, prepared in 10% ethanol, 90% sunflower oil) was administered via modified gavage once a day for 3 d. Animals were allowed to recover for two weeks, and dosed again for 3 d to ensure complete induction of cre expression. This protocol has been well established as sufficient to induce recombinase activity in SLICK animals (Wilhelm et al., 2012; Zhu et al., 2016). There was a minimum of two weeks between treatment and retrograde tracer injections in SLICK animals. In Cx3cr1<sup>creER</sup>+ animals, this interval was extended to four weeks to ensure specificity of gene deletions within only microglia, as described above.

Retrograde tracer injections

This study focused on one motor pool axotomized after sciatic nerve injuries, the motoneurons innervating the lateral gastrocnemius (LG) muscle. LG Motoneurons were retrogradely labeled either by intramuscular injection of the long-lasting, non-toxic tracer fast blue (FB; Polysciences, Inc) or an adeno-associated virus expressing mCherry (AAV1-mCherry).

FB was injected unilaterally or bilaterally into the LG muscles of adult animals. The animals were anesthetized with isoflurane (4% induction, 2% maintenance) and given preoperative buprenorphine (0.05 mg/kg, i.p.). Once the animals reached a surgical plane of anesthesia, a small skin incision was made to expose the LG muscles, and a 10-μL Hamilton syringe was introduced to inject 2–5 μL of a 1.5% solution into the belly of the muscle. Skin incisions were sutured and the animals were allowed to recover for at least 7 d before nerve surgery to ensure complete retrograde transport.
To visualize the full dendritic arbor of axotomized motoneurons, a small subset of animals were injected with AAV1-mCherry (2 μl of 10²⁵ IU/mL) into the LG muscle at postnatal day 15 with the same anesthesia and analgesia regime as above. Viral injections were performed with glass microelectrodes inserted into the muscle through the skin. These animals were allowed to survive until adulthood (two to three months) before further manipulation. Injections in young animals greatly increase AAV1 infection rates.

**Surgeries**

All nerve surgeries were performed under isofluorane anesthesia (4% induction, 2% maintenance) and the animals were given preoperative 0.05 mg/kg buprenorphine (intra-peritoneal) to manage possible post-operative pain. The spinal nerves were transected at the mid-thigh skin incision to expose the hindlimb muscles, which includes the axons of the LG and other spinal motoneurons. A small cutaneous flap was reflected and the overlying biceps femoris. Intrinsic hindlimb muscles were exposed by a mid-thigh skin incision and blunt dissection of the overlying biceps femoris. In animals in the cut/ligated condition, a silk suture was tied around the sciatic nerve where it was secured with fibrin glue (SILASTIC film (Dow Corning No. 501-1) was placed beneath the sciatic nerve). A small rectangle of PARAFilm (M Medical Products) was placed beneath the sciatic nerve, which includes the axons of the LG and other spinal motoneurons, a small subset of animals were injected with AAV1-mCherry (2 μl of 10²⁵ IU/mL) into the LG muscle at postnatal day 15 with the same anesthesia and analgesia regime as above. Viral injections were performed with glass microelectrodes inserted into the muscle through the skin. These animals were allowed to survive until adulthood (two to three months) before further manipulation. Injections in young animals greatly increase AAV1 infection rates.

**Table 2. Primary antibodies used for immunohistochemistry (IHC) and Western blottings (WB)**

| Antigen        | Immunogen                                      | Host/type          | Manufacturer                  | RRID #        | dilution | Use     |
|----------------|------------------------------------------------|--------------------|--------------------------------|---------------|----------|---------|
| KCC2           | N-terminal His-tag fusion protein; pan-rat KCC2; aa 932–1043 | Rabbit/ polyclonal | Millipore catalog #07-432 | AB_11213615   | 1:500    | IHC     |
| NeuN           | Purified cell nuclei from mouse brain A60 clone | Mouse/ monoclonal  | EMD Millipore catalog #MAB377 | AB_2298772    | 1:500    | IHC     |
| EGF            | Recombinant GFP 6-his tag | Chickens/ polyclonal | Serotec catalog #601644 | AB_10000240   | 1:1000   | IHC     |
| Iba1           | C terminus of Iba1 | Goat/ polyclonal | Novus catalog #NB 100-1028 | AB_521594     | 1:500    | IHC     |
| mCherry        | Purified recombinant peptide produced in E. coli | Goat/ polyclonal | MyBioSource catalog #MBS448050 | NA            | 1:100    | IHC     |
| NF-H           | Purified NF-H from bovine brain                | Chicken/ polyclonal | Aves catalog #NFH | NA            | 1:500    | IHC     |
| VACHT          | Recombinant protein; rat VACHT; aa 475–530 | Guinea pig/ polyclonal | Synaptic Systems catalog #13906 | NA            | 1:500    | IHC     |
| ChAT           | Human placental enzyme | Goat/ polyclonal | Millipore catalog #AB144P | AB_2079751    | 1:100    | IHC     |
| ATF3           | Recombinant protein corresponding to aa MMLDHPGVSASEVSASAIVPCLSSPGSLVLEDANLFVKERLFAINGQVHLCHRNSSALESVTVSDRPLGVSKVAPAEEEDRKKRRRERKIAAACRINKK| Mouse/ polyclonal | Novus NBP2-34489 | NA            | 1:200    | IHC     |
| MAPK (Erk1/2)  | Synthetic phosphopeptide; human p44 | Rabbit/ polyclonal | Cell Signaling Technology catalog #9101 | AB_331646     | 1:1000   | WB      |
| β-actin        | Synthetic phosphopeptide; human β-actin; N-terminal region | Rabbit/ polyclonal | Novus catalog #NB600-503 | NA            | 1:5000   | WB      |

Most sections were reacted with primary antibodies against KCC2 and NeuN, and some sections additionally included antibodies to detect microglia (anti-Iba1 or anti-GFP for CX3CR1-GFP amplification in mice carrying this reporter protein). To investigate potential differences between γ and α motoneurons, these sections were reacted with an antibody against ChAT and ATF3 to confirm the injected motoneuron population (Table 2). ATF3 is upregulated in the nucleus of all axotomized neurons (Tsujino et al., 2000). In all cases, the primary antibody against KCC2 was raised against a peptide sequence containing aa 932–1043, which is shared by both KCC2 isoforms (KCC2a and KCC2b). Sections were then washed in PBST and incubated with appropriate fluorescent-coupled species-specific secondary antibodies for 2 h (1:100 in PBST; Table 3). Immunoreactivity (IR) to KCC2 was visualized after incubating with an Alexa Fluor 647-coupled secondary antibody, NeuN-IR with cyanine Cy3-coupled secondary antibodies. LG motoneurons retrogradely labeled with FB were visualized by their fluorescence (excitation 405 nm; emission 420 nm). After washing, the sections were followed by paraformaldehyde (PFA; 4% in 0.1 M phosphate buffer). The spinal cord and injected muscles were harvested and postfixed overnight in 4% PFA before cryoprotection in 30% sucrose for at least 24 h. The spinal cord dura was then removed and the L3–L5 segments isolated. The spinal cords were sectioned in a transverse plane on a freezing sliding microtome at 50-μm thickness and collected free-floating. To improve antibody penetration, the sections were heated for 20 min in 0.01 M sodium citrate with 0.05% Tween 40 (pH 6, maximum temperature 95°C), washed in 0.01 M PBS with 0.3% Triton (PBST), and blocked in normal donkey serum (NDS; 10% PBS with 1% Triton) for 1 h. The tissue was then incubated, while shaking for two nights at room temperature, with a mixture of primary antibodies diluted in PBST. The antibodies used, their sources, and RRID numbers are listed in Table 2.

**Tissue collection, processing, and immunocytochemistry**

Animals were allowed to survive 3, 7, 14, 21, 28, or 60 days (d) after nerve surgeries, depending on the experiment. They were then overdosed with Euthanol (100 mg/kg) and transcardially perfused with saline-heparin followed by paraformaldehyde (PFA; 4% in 0.1 M phosphate buffer). The spinal cord and injected muscles were harvested and postfixed overnight in 4% PFA before cryoprotection in 30% sucrose for at least 24 h. The spinal cord dura was then removed and the L3–L5 segments isolated. The spinal cords were sectioned in a transverse plane on a freezing sliding microtome at 50-μm thickness and collected free-floating. To improve antibody penetration, the sections were heated for 20 min in 0.01 M sodium citrate with 0.05% Tween 40 (pH 6, maximum temperature 95°C), washed in 0.01 M PBS with 0.3% Triton (PBST), and blocked in normal donkey serum (NDS; 10% PBS with 1% Triton) for 1 h. The tissue was then incubated, while shaking for two nights at room temperature, with a mixture of appropriate primary antibodies diluted in PBST. The antibodies used, their sources, and RRID numbers are listed in Table 2.

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Sections containing AAV1-mCherry motoneurons were reacted with antibodies against KCC2 and IR was detected with Cy5-conjugated secondary antibodies. In these sections, the mCherry signal was amplified by incubating with an antibody against mCherry revealed with Cy3-conjugated secondary antibodies. In addition, these animals were CX3CR1-GFP, and GFP was amplified with anti-GFP and revealed with FITC-conjugated secondary antibodies.

In muscles harvested from sciatic nerve transection-repair animals, cryostat sections were cut at 25-µm thickness and collected onto glass slides. After washing, the slides were incubated for 1 h with 10% NDS. Then they were incubated overnight at room temperature with α-bungarotoxin conjugated to Alexa Fluor 555 (αBgTx-555, Invitrogen catalog #B13422, 1:100) and primary antibodies against neurofilament-H (NF-H) to label the axonal cytoskeleton in thick motor axons and against the VACHT to label the presynaptic motor nerve terminals (Table 2). VACHT-IR was revealed with Cy5-conjugated secondary antibodies and NF-H with FITC-conjugated antibodies (dilution 1:100; Table 3). Sections were mounted with VECTASHIELD and 50 motor endplates per animal were imaged for quantification with an Olympus FV1000 Confocal microscope (20×).

Quantification of KCC2-IR in the cell surface

Motoneurons in which somatic KCC2 levels were measured were always filled with FB and imaged at 60×1. Unless indicated otherwise, 10 FB+ motoneurons ipsilateral to the injury were analyzed. In the case of animals with the contralateral side used as intact control, 10 FB+ motoneurons were also analyzed on the contralateral side. For all motoneurons, a single z-plane (focal depth ~0.49 µm at 633-nm excitation with refractory index n = 1.518 oil immersion 60× objective of NA 1.35) was selected for each FB labeled motoneuron studied. These planes always included a nucleolus and had minimal dendritic departures from the soma. NeuN-IR was used to aid in finding mid-plane optical sections within the z-stack. KCC2 immunofluorescence intensity around the soma of each motoneuron was quantified using FIJI (Fig. 1C). The Wand (tracing) Tool (mode: 8-connected; tolerance: 1000) was used to automatically select the edge of the labeled soma containing FB. This tracing was converted to a 0.42-µm wide line superimposed on the neuronal plasma membrane. The average KCC2 Cy5 fluorescent intensity along the line was calculated (gray level, 12 bits). In each section, background measurements were taken from a 9 µm² square region of the neuropil in the same optical plane, adjacent to the motoneurons and lacking labeled dendrites. KCC2 immunofluorescence was corrected against this background level by calculating the percentage higher than background [100 × (membrane intensity average – background intensity average)/average background intensity]. Data points are presented as averages for each animal, each obtained from 10 motoneurons per animal and side of the spinal cord. Different conditions (i.e., injury, sham, time after injury, genotype, spinal cord side) were compared by obtaining averages of n = 4–7 animals.

To investigate the differences between α and γ motoneurons, motoneurons in Lamina IX were visualized with ChAT and those axiomatized by sciatic transection were identified with ATP3 labeling. Injured motoneurons and those in the same motor pool on the contralateral side were manually traced because ChAT is downregulated after injury and cannot be reliably recognized automatically. Quantification of KCC2-IR was performed as described above. The neurons were designated as γ motoneurons if they had mid-plane cross sectional areas smaller than 475 µm² (criteria published in Shneider et al., 2009) and equivalent numbers of injured and uninjured α and γ motoneurons were quantified (n = 16 injured, n = 25 uninjured).

Quantification of KCC2-IR in the cytoplasm

In animals euthanized 3 d after sciatic nerve cut and ligation, KCC2-IR was quantified in the cytoplasm to address whether KCC2 disappearance from the membrane may have been, in part, explained by increased KCC2 internalization or decreased insertion into the membrane. In the same 10 motoneurons quantified for membrane KCC2-IR, samples of KCC2-IR were taken from five 9-µm² rectangles per motoneuron placed on the cytoplasm. These areas were selected in the FB channel to eliminate the potential for bias and they did not include areas of the nucleus or nucleolus. The average cytoplasmic KCC2-IR was calculated for each motoneuron and then corrected for background as described for the membrane measurements.
Qualitative analyses of KCC2 in motoneuron dendritic arbors

Spinal cord sections containing AAV1-mCherry filled motoneurons and KCC2-IR were imaged at high magnification (60x1). At this magnification, the field of view includes only a small region of the whole dendritic arbor in the section. To image the entirety of the dendritic arbor contained within the section we used image tiling of contiguous z-stacks. The images were imported into Neurolucida (v10.0, MBF Bioscience) to fully reconstruct the cell bodies and dendritic arbors contained within the 50-μm-thick section. The dendritic arbors contained within the section were reconstructed in seven motoneurons 14 d after axotomy. Following reconstruction, KCC2-IR labeling in dendritic processes was designated as complete KCC2 depletion (Fig. 2, green), partial depletion (Fig. 2, blue), or normal levels of KCC2 (Fig. 2, pink), in comparison to relative levels of KCC2 in dendrites in the surrounding neuropil. The distances of these dendritic segments to the cell body and their total surface were tabulated, and a Sholl analysis was performed to calculate the proportion of membrane covered with KCC2-IR at different distances from the cell body, using 100-μm bins.

ANA-12 treatment

A subset of WT animals was treated continuously from injury to euthanasia with ANA-12, a competitive TrkB antagonist, or vehicle. ANA-12 was prepared in DMSO (30%), PEG300 (30%), and sterile saline (40%) at 2 mg/ml, and delivered (0.5 μl/h) intraperitoneally using an ALZET-2002 mini-osmotic pump. This rate was chosen to deliver the same concentration of ANA-12 per 12 h as when administered twice daily with intraperitoneal injections (Ambrogini et al., 2013; Chen et al., 2015). The use of osmotic pumps allows maintenance of a more consistent concentration of the drug throughout the treatment period. Pumps were filled with ANA-12 or vehicle and primed in sterile saline overnight at 37°C. Pumps were implanted in the animals at the time of nerve cut/ligation (ANA-12, n = 6; vehicle, n = 4) or sham (ANA-12, n = 4). They were brought from warm saline and immediately inserted into the peritoneal cavity, being careful not to disturb internal organs. The wound was closed in layers and animals were monitored daily to insure pump stability. Before perfusion 14 d later, pumps were removed, weighed, and evacuated with a needle to ensure the drug was properly delivered. All pumps had expelled the expected volume of drug.

A subset of animals (n = 2 per group: sham-vehicle, sham-ANA-12, injury-vehicle, injury-ANA-12) was used to confirm the effectiveness of the ANA-12 treatment. Western blots were used to analyze phosphorylation of ERKs (pERKs), a downstream effector of TrkB activation known to increase in the spinal cord after nerve injury (Wang et al., 2004). Animals were treated with ANA-12 or vehicle (as above) and underwent bilateral sham or sciatic nerve cut/ligations. Seven days postinjury, the animals were euthanized and rapidly perfused with ice-cold saline, and the lumbar spinal cords were extracted and flash frozen. The tissue was homogenized in phosphatase-inhibitory lysis buffer (10 mM HEPES, pH 7.4; 50 mM NaF; 50 mM NaCl; 5 mM EDTA; 5 mM EGTA; 0.1% Triton X-100; 1 mM Na3VO4; 10 μM leupeptin; and 1 mM phenylmethylsulfonyl fluoride, 25 kIU/ml aprotinin) and centrifuged (13,000 x g at 4°C) to separate lysate. The pellet was discarded and cell lysates preserved at –80°C until the samples
were assayed for protein concentration and processed for PAGE. Protein concentration was determined with the Bio-Rad DC protein assay (catalog #5000116) at 750 nm. Protein standard dilutions were prepared in buffer containing the protease inhibitors (see above). Forty micrograms of protein from each sample was loaded onto Bio-Rad 10% polyacrylamide precast gels (50 μl wells, catalog #456-0834) and Bio-Rad Kaleidoscope protein

Figure 2. KCC2 protein is lost extensively from axotomized motoneuron dendrites 14 d after sciatic cut/ligation. A, AAV1-mCherry filled LG motoneurons (A1) and KCC2-IR in the same section (A2) 14 d following sciatic cut/ligation. Note some afferents are also labeled in the dorsal horn after AAV1 mCherry muscle injections. Scale bar = 200 μm. B, High-magnification confocal image through the soma and proximal dendrites of two mCherry-labeled and axotomized motoneurons (mCherry-red). KCC2 immunofluorescence (white) was not observed around the cell body and the proximal dendrites. Scale bar = 50 μm. C, The full dendritic arbor was imaged at high magnification using confocal image tiling and then the neurons reconstructed in 3D using Neurolucida. Example of six individual motoneurons (color-coded) reconstructed within one section with KCC2-IR in the background. Note the extension of dendrites toward the dorsal horn and also entering the white matter. Scale bar = 200 μm. D, Dendrites were color-coded to indicate complete KCC2 loss (green, “none”), partial KCC2 loss (blue), or normal KCC2 levels (pink), relative to surrounding dendrites in the neuropil. Loss of fluorescence in the proximal dendrites was similar to that on the soma membranes (green and region i). IR to KCC2 was partially reduced (blue and region ii) along dendrites for various distances. However, no significant loss of KCC2 was found on distal dendrites; scale bars (i, ii) = 5 μm. E, Box plots of the percentage of surface area in each category (complete depletion, partial depletion, or no depletion) for all dendrites for which the complete length was traceable (not cut short during sectioning). More than half of the available dendritic surface area was completely depleted (Table 4; *** p < 0.001). Box plots represent 25th, median, and 75th percentiles; whiskers = 10th and 90th percentiles. Crosses = mean; points represent average for individual neurons (n = 6, 3.12 ± 0.98 dendrites/animal). F, G, Sholl analysis (100-μm bins) of dendritic length and KCC2 depletion. F, Average (± SD) percentage of dendritic lengths within each bin (n = 7 neurons, 5.29 ± 1.1 dendrites per neuron), illustrating that within the first 200 μm, the majority of dendrites have no KCC2 visible, whereas normal KCC2 levels can be observed at the most distal ends of dendrites. G, Isolated single motoneuron from C separated for Sholl analysis. The majority of KCC2 is completely lost in the first 200 μm of dendrite; at 400 μm, dendritic KCC2 is largely preserved. H, Example dendrogram of the motoneuron in G, color-coded for KCC2 depletion. Dendrite branches ending in circles represent dendrites that were fully contained within the section (quantified in E). Dendrite branches that were cut during sectioning and could not be fully traced are designated with slashes. The majority of dendrites for which the ending could be observed had normal levels of KCC2 present in their terminal regions.

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molecular weight markers (catalog #161-0375) were loaded into at least one well on each gel. After electrophoresis, Western blotting was conducted overnight at 4°C onto PVDF membranes (Bio-Rad catalog #162-0177) using 150-ma constant current. Primary rabbit antibodies against phospho-p44/42 MAPK (pMAPK) or pERK (Thr202/Thr204) and β-actin were used (Table 2). The secondary antibody in all cases was HRP-conjugated donkey anti-rabbit, and protein immunostaining was revealed using chemiluminescence (Clarity ECL Western substrate, Bio-Rad catalog #170-5061). β-Actin was used as a loading control. The membrane was stripped using Re-Blot Plus, Strong Solution (Millipore catalog #2504) before re-probing with anti β-actin.

Pharmacogenetic TrkB inactivation

F616A mice carry a mutation in the TrkB receptor that makes it sensitive to specific inhibition by the small molecule inhibitor 1-(1,1-dimethylethyl)-3-(1-naphthalenylmethyl)-1H-pyrazolo[3,4-d]pyrimidin-4-amine, PP1 analog (1NMPP1). We administered 1NMPP1 following protocols well established in the literature (Chen et al., 2005). Briefly, 3 d before nerve injury, F616A animals were switched from normal drinking water to drinking water with 1NMPP1 (5 μM, 0.01% DMSO) or vehicle. Animals were monitored twice daily to ensure proper hydration and adequate access to the water.

KCC2 mRNA detection using RNA-Scope

Standard surgical and euthanasia procedures were performed as described above with the exception that perfusions were performed with 4% PFA in 1× PBS. For RNA-Scope labeling, we prepared animals with sciatic nerve injury ligation. Animals were euthanized 3, 7, and 14 d after nerve surgeries (n = 3 animals per group). Following perfusion, spinal cords were harvested and post-fixed overnight at 4°C, followed by stepwise cryo-protection (10% sucrose in 1× PBS overnight, 20% sucrose in 1× PBS overnight, 30% sucrose in 1× PBS overnight) before sectioning (16 μm) on a cryostat. Sections were collected on RNase free slides. Samples were then sent to Advanced Cell Diagnostics (ACD Bio) on dry ice and processed for RNA-Scope KCC2 mRNA in situ hybridization using a probe directed toward the 501- to 1717-bp region of the KCC2 mRNA (Mm-SLC12A5 ACD catalog #311901). This probe detects mRNA for both KCC2a and KCC2b isoforms and consists of 20 Z probe pairs. We used the ubiquitously expressed peptidylprolyl isomerase B (cyclophilin B; ACD catalog #313911) as positive control to confirm mRNA preservation, and dihydrolipoic reductase (dapB), a gene from Bacillus subtilis (ACD catalog #310043), as negative control to assess background labeling. To improve tissue adherence, sections were baked at 60°C for 45 min and then further postfixed on the slide (4% PFA in 1× PBS) for 90 min at room temperature before serial dehydration (5 min each; 50% EtOH, 70% EtOH, 100% EtOH, 100% ETOH) and air-drying. Standard RNA-Scope procedures (Wang et al., 2012) were used and the KCC2 mRNA binding detected with RNA-Scope 2.5 HD Reagent Kit-BROWN (catalog #322300). The sections were counterstained with standard Nissl stain (hematoxylin).

RNA-Scope KCC2 mRNA image analysis

Dark- and bright-field images were taken on an Olympus BX60 microscope with Spotcam RX3 camera at 10× and 40× 2 magnification, respectively. Images were imported into ImagePro ver 7.0 (Media Cybernetics), and thresholds for KCC2 mRNA signal were determined to segment the KCC2 mRNA labeling from the image. The same thresholds were used for all sections, but different thresholds were used for the cytoplasm and nucleus as the Nissl stain had a different gray level in nuclei and most KCC2 mRNA signal was usually concentrated on top of the nucleus. The total area of cytoplasmic or nuclear labeling was calculated (Fig. 3D–F) and data presented as average percentage coverage of KCC2 mRNA labeling in each cell compartment. We noted interanimal variability in the strength of the KCC2 mRNA signals. For this reason, we always compared labeling intensities to the contralateral control uninjured side.

Statistical analysis

Statistical tests were performed as described in the results and powered to power > 0.8 for α < 0.05 (see statistical tables; Tables 4–6). Data were evaluated with t tests or one-way or two-way ANOVA as appropriate. Bonferroni post hoc multiple comparisons tests were performed against controls (WT or transgene negative control animals) unless otherwise described. Finally, RNA-Scope quantification was compared using paired t tests between spinal cord sides ipsilateral and contralateral to the injury. Power was calculated using SigmaPlot (Systat Software, Inc., 12.3) and all other statistics were performed with PRISM (GraphPad, 7.03).

Results

KCC2 is lost from the somatic and proximal dendrite membrane of motoneurons axotomized after nerve injury

Using antibodies raised against a rat KCC2 sequence (aa 932–1043) shared by both KCC2 isoforms (KCC2a and KCC2b), we found a loss of KCC2-IR on the sciatic motor pools ipsilateral to the injury after unilateral sciatric nerve transections (Fig. 1). Fourteen days after sciatric nerve transection, KCC2-IR was strongly reduced or absent from the plasma membranes of cell bodies and proximal dendrites of LG motoneurons labeled with FB (Fig. 1A,B). Using an unbiased automatic detection of the cell body plasma membrane based on FB fluorescence (see Materials and Methods; Fig. 1C), we measured KCC2-IR along the membrane. This measurement was then normalized to background fluorescence estimated in the same optical plane. KCC2-IR levels are expressed as a percentage of fluorescence intensity above background (0% = background). Ten motoneurons were studied in each animal, unless otherwise indicated. Thus, n always refers to the number of animals analyzed.

In ~90% of the motoneurons analyzed in sham or naïve animals, levels of KCC2 fluorescence averaged between 75% and 150% above background (Fig. 1D). After injury, average KCC2 immunofluorescence estimates dropped to around 50% higher than background, with more than
half of the estimates below this level (Fig. 1D). At this level of measured KCC2 immunofluorescence above background, no KCC2 immunofluorescence was visible along the plasma membrane. This level above background might be due to an offset gray level imposed by the cytoplasm that is always slightly more fluorescent than neuropil background. Similar numbers of male and female mice underwent axotomy, and at 14 d after injury a significant depletion was found in injured animals, but no sex differences were detected in either intact or injured motoneurons (Fig. 1D). A two-away ANOVA for condition (injured vs non-injured) and sex showed significant differences according to condition, but not sex, or the interaction of injury and sex (Table 4). Thus, males and females were pooled together in all experiments described below.

Despite the robust loss of KCC2 on the soma, KCC2-IR can be observed on some small dendrite segments within the neuropil around the cell bodies (Fig. 1A, B, yellow

Figure 3. KCC2 mRNA is lost by 3 d after peripheral axotomy. A, Darkfield image of a lumbar 4 section 3 d after unilateral sciatic nerve cut/ligation, processed using RNA-Scope. KCC2 mRNA visualized in white. Scale bar = 200 μm. Very little KCC2 mRNA is found in the area around the injured motor pool (right) in Lamina IX (LIX), relative to the contralateral intact motor pools. B, C, Brightfield images of motoneurons on the intact (B) or cut/ligated (C) sides of the spinal cord 3 d after PNI. Injured motoneurons lose KCC2 mRNA, whereas motoneurons contralateral to injury and interneurons (INs) or γ motoneurons (γMNs) on both sides of the spinal cord maintain KCC2 mRNA. Nuclei from glia are also visible and are more densely concentrated around axotomized neurons. KCC2 mRNA can also be seen extending into dendritic processes of intact motoneurons (rectangle in B, inset). Scale bar = 25 μm. D–F, RNA-Scope quantification procedure. Thresholding was used to highlight KCC2 mRNA (red). Nissl staining caused differences in background intensity, so difference thresholds were required for cytoplasm and nuclear quantification. Thus, the outlines of the motoneuron somata were traced (green) with the nucleus excluded to get the area of KCC2 mRNA within the cytoplasm (D). Nuclei (E) were traced and threshold separately. The total area, including cytoplasm and nucleus (F), was also traced for comparison of total area to KCC2 mRNA area. G, H, Percentage KCC2 coverage of soma (G) and nucleus (H). Animals were euthanized 3, 7, or 14 d (n = 3/group) after unilateral sciatic nerve cut/ligation, and sciatic motoneurons were quantified as described above. There were no differences between levels of KCC2 between time points; only injury state predicted KCC2 levels (Table 4). Thus, data from all time points were pooled and paired t tests were performed comparing motoneurons ipsilateral and contralateral to injury (Table 4; ***p < 0.001).
AAV1-mCherry into the LG. AAV1-mCherry fills the dendritic arbor, P15 animals were injected with soma and most proximal dendrites; therefore, to visualize plasma membrane 14 d after nerve injury. FB only fills the responses to GABA/glycine. Thus, we performed experiments, minimizing potential chloride accumulation and chloride from somatic and proximal dendrite compartments, observing on dendrites it could act as a siphon to extrude motoneurons. We were concerned that if KCC2 was preserved in sciatic motoneurons or that they originate from controls. It is possible they are distal segments of axotomized sciatic motoneurons (Fig. 2A,E). These animals underwent sciatic cut/ligation in adulthood, and we analyzed seven motoneurons 14 d after injury. Three-dimensional reconstructions of motoneuron dendritic arbors were obtained using NeuroLucida (Fig. 2C,D). In the proximal dendrites, KCC2 was depleted to a similar extent as in the soma. However, we observed a gradient in which more KCC2-IR could be observed on the dendrites at progressively greater distances from the soma, and on the most distal dendrites KCC2-IR was fully preserved. The extent of KCC2 depletion varied from neuron to neuron and dendrite to dendrite. Overall, 52.6 ± 13.9% (± SD) of the total dendritic surface area had no KCC2-IR, 33.0 ± 8.3% showed a

| A | 1D | Non-normal distribution, unequal variance | Two-way ANOVA | Power and statistical significance |
|---|---|---|---|---|
| B | 2E | Normal distribution, unequal variance | One-way ANOVA | Power = 0.999 |
| C | 3H | Normal distribution, unequal variance | Two-way ANOVA | Power = 1.0 |
| D | 3G | Normal distribution, unequal variance | Two-way ANOVA | Power = 0.981 |
| E | Sham vs ligated | One-tailed t test | Power = 0.95 |

Table 4. Details of statistics for experiments describing the phenomenon of KCC2 loss on spinal motoneurons following axotomy

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partial preservation of KCC2-IR, and 14.3 ± 10.8% of dendrites surface showed normal KCC2-IR 14 d after injury (Fig. 2E). The differences in total surface area with no, partial, or normal KCC2-IR were statistically significant (Table 4). Using Sholl analysis, we found that the areas with full preservation of KCC2 were always distal and regions with total depletion were always proximal (Fig. 2F). In dendritic segments located in the first 100 μm of Sholl distance from the cell body, 87.3 ± 14.9% of their surface was deplet ed of KCC2, and at 100- to 200-μm distance 47.4 ± 26.8% of the surface was totally depleted. However, in dendritic segments above 200 μm of Sholl distance, partial or total preservation of KCC2-IR was commonly observed (Fig. 2F–H). In analyses of individual dendrites (Fig. 2G, H) complete preservation of KCC2-IR was found in the terminal dendritic regions (Fig. 2, dot marks in dendrograms). Not all dendrites were fully contained within the section (hatch marks in dendrograms) and these were not included in the Sholl analysis; mCherry-labeled dendrites ending within the top and bottom surfaces of the section typically display a few ending varicosities. KCC2 preservation in the most distal dendrites was independent of whether the dendrites ended in the gray or white matter (Fig. 2C). The impact of KCC2 retention in these distal regions on overall chloride levels might be minimal since motoneuron dendrites at these distances are of very small caliber and contribute little to the total surface; thus, the overall percentage of dendrite surface with normal KCC2 density is quite small (Fig. 2E).

On the other hand, the impact of partial retention of KCC2 density in mid-distal dendrite regions is yet unknown (see Discussion).

KCC2 regulation on axotomized motoneurons occurs at the transcriptional level

Disappearance of KCC2 protein from the membrane of axotomized motoneurons could occur because of membrane protein turnover and degradation, slower KCC2 protein trafficking to the membrane, or reduced gene expression. In dorsal horn interneurons and spinal motoneurons after SCI, KCC2 downregulation has been reported to occur rapidly and through post-translational KCC2 phosphorylation mechanisms likely affecting trafficking and membrane stabilization of KCC2 (Coull et al., 2005; Boullenguez et al., 2010; Bos et al., 2013; Ferrini and De Koninck, 2013). However, previous studies in axotomized brainstem cranial motoneurons reported a strong downregulation of KCC2 mRNA after PNI. This results in slower removal of KCC2 from the membrane, but the loss is more profound and longer lasting (Toyoda et al., 2003).

To investigate whether the loss of KCC2 after PNI is the result of reduced gene expression, we measured kcc2 mRNAs using a diaminobenzidine (DAB)-based RNA-Scope method and a probe that recognizes a 501- to 1717-bp region shared by both kcc2 isoforms (Fig. 3). In dark-field microscopy we observed complete depletion of kcc2 mRNA in Lamina IX motoneurons in the region corresponding to the sciatic motor pool ipsilateral to the lesion and little changes in other spinal cord laminae or in Lamina IX motoneurons on the intact contralateral side (Fig. 3A). At high magnification, there was robust and consistent kcc2 mRNA signal in the cell bodies and nucleus of all uninjured spinal cord Nissl-stained neurons (Fig. 3B). Within the nucleus of these cells, there was frequently a large spot of reaction product localized on the nucleolar region. In motoneurons (recognized by their large size in Lamina IX) we also observed kcc2 mRNA reaction product entering proximal dendrites (Fig. 3B). Both cytoplasmic and nuclear labeling were strongly decreased in axotomized motoneurons (Fig. 3C). We quantified RNA-Scope reaction products by measuring the area covered by reaction particles within both the cell body cytoplasm and nucleus of motoneurons in Lamina IX sciatic motor pools. Different threshold values were used in each cellular compartment (see Materials and Methods) because of differences in intensity of the reaction product and the different gray levels of the Nissl-stained nucleus versus cytoplasm (Fig. 3D–F). Since some interanimal variability was observed in the strength of the RNA-Scope reaction, we compared motoneurons ipsilateral and contralateral to the injury within animals. By 3 d after injury, the extent of coverage by kcc2 mRNA reaction product was reduced by 90.7 ± 1.5% (SE) in the cytoplasm and 89.8 ± 0.6% in the nucleus. These depletions were consistently maintained through 14 d after injury (81.7 ± 2.3% and 80.1 ± 9.4% depletions in cytoplasm and nucleus, respectively). There were no significant differences in the percentage depletion of the kcc2 mRNA signal in cytoplasm or nucleus 3, 7, and 14 d after injury (Table 4), thus data from all three time points were pooled together. Paired t tests showed highly significant reductions in kcc2 mRNA signal in both cytoplasm and nucleus.

Because KCC2 mRNA was depleted by 3 d after injury, and our time course of KCC2 protein depletion showed a trend toward KCC2 loss on the soma at this time point, we further investigated this time point by examining KCC2 distribution on AAV1-mCherry labeled motoneuron soma and dendrites of animals 3 d after sciatic nerve cut and ligation. Qualitatively, these motoneurons showed no large decreases in KCC2-IR on the soma or throughout the dendrites, further confirming that the mRNA disappears before differences in protein can be observed (Fig. 4). This leads to suggesting that KCC2 loss is due to depletions in kcc2 transcription and turn-over of membrane KCC2 without replacement by new protein. The possibility that concurrent increased internalization of KCC2 protein is also contributing to the loss observed was tested by quantifying KCC2-IR in the cytoplasm of motoneurons 3 d postinjury. We found no significant differences (t test; Table 4) between cytoplasmic KCC2-IR in motoneurons of sham animals (45 ± 7.9% higher than background) and animals in which the sciatic nerve was cut and ligated 3 d prior (50 ± 23.5%). While this finding suggests that the initial depletion of KCC2 is not due to increased internalization of KCC2, it is still possible that any internalized protein is quickly degraded and undetectable with immunocytochemistry. The rate of KCC2 recycling in intact or injured motoneurons is unknown but the rate of KCC2 loss seem quite slow for any significant contributions from rapid enhanced internalization.
RNA-Scope preparations revealed that kcc2 mRNA was preserved in some small neurons near the large motoneurons we quantified. These could be interneurons or small motoneurons. It is impossible to definitively distinguish these different types of neurons in our Nissl-stained RNA-Scope preparations, so we investigated potential differences between γ and α motoneuron KCC2 expression and dynamics by labeling all motoneurons with ChAT antibodies and distinguishing each type based on cell size. γ Motoneurons were not reliably filled by retrograde tracers, so overlapping ChAT and ATF3-IR was used to determine the injured motoneurons. As reported elsewhere, injured motoneurons downregulated ChAT (Ichimiya et al., 2013) and injured motoneuron nuclei were clearly visible by ATF3 labeling (Fig. 5A, B). Small motoneurons (<475-μm² cross sectional area) were designated as γs, as previously defined (Shneider et al., 2009). KCC2-IR was quantified on the somatic membrane, and although no significant differences between intact α and γ motoneurons (n = 25 per group) were detected there was a trend toward lower KCC2-IR in γs. Injured γ and α motoneurons showed significant lower KCC2-IR relative to intact α motoneurons (Table 4). However, KCC2 in γ motoneurons was more variable and many showed KCC2 at normal levels (Fig. 5C). This variability and the trend toward diminished KCC2 in γs resulted in no significant differences between intact and uninjured γ motoneurons. Thus, it is possible that γ motoneurons lose KCC2 with different time-courses or not at all, or that there are differences according to the type of γ motoneuron (dynamic vs static). We focused all our analyses on the larger population of α motoneurons that reliably lose KCC2 by 14 d. The interesting possibility of significant differences between γ and α motoneuron KCC2 expression and reactions to injury will need to be investigated in future work.

**Time course of KCC2 protein loss and recovery and its relation with microglial reactivity and muscle innervation**

To investigate the time course of KCC2 protein down-regulation on axotomized motoneurons and a possible relationship with activation of ventral horn microglia around injured motoneurons, we performed nerve injuries in male and female Cx3cr1GFP/− heterozygous mice in which microglia express GFP. To analyze the effect of regeneration and muscle innervation in KCC2 expression, we either prevented regeneration by cutting and ligating the sciatic nerve (tying the proximal stump with silk) or enabled regeneration by cutting and repairing the nerve (end-to-end anastomosis of the proximal and distal nerve stumps using fibrin glue). Analyses of KCC2 immunofluorescence were performed in cohorts of animals either without a nerve injury (intact) or 3, 14, 21, 28, or 60 d after injury (n = 4 or 5 per group). All analyses were performed on the cell bodies of 10 MNs per animal (as in Fig. 1). Animals with sciatic nerve repair were only analyzed at 21, 28, and 60 d for comparison with cut/ligated animals, as this is when differences in regeneration and muscle reinnervation occur between injury models.
At low magnification, KCC2-IR was diminished in the Lamina IX region containing injured sciatic motoneurons (including the FB retrogradely labeled LG motor pool). This area closely overlapped with the ventral horn region showing a strong microglial reaction (Fig. 6A). Based on KCC2 quantification on the cell body membrane of FB-labeled LG motoneurons, significant differences were found between control and axotomized motoneurons at different times after injury with regeneration allowed or prevented (Table 4). Using post hoc comparisons (Bonferroni-corrected t tests), we found that membrane KCC2-IR was significantly greater in control motoneurons than almost all other groups. Notably, KCC2 levels on the somatic membrane of repaired motoneurons 60 d after injury were similar to controls in average and variance (control 125.4 ± 48.4% above background vs 120.9 ± 33.9% in regenerated motoneurons 60 d after injury).

Similar regulation of KCC2 after axotomy was observed in WT animals and Cx3cr1GFP/+ heterozygous animals (see below) as well as Cx3cr1KO/GFP animals. We observed no changes in either microgliosis or KCC2 down-regulation in Cx3cr1GFP/GFP animals, where both alleles of the fractalkine receptor are replaced by GFP and thus are Cx3cr1KOs (data not shown). KCC2 regulation on the cell bodies of axotomized motoneurons is thus independent of CX3CR1 function. Moreover, microgliosis in the ventral horn is largely resolved by 60 d after injury even when regeneration is not allowed (cut and ligated sciatic nerves; Fig. 6C,D), without recovery of KCC2.

One interpretation of the above results is that KCC2 restoration on the membranes of motoneurons after PNI occurs only following regeneration of their axons. To investigate whether KCC2 restoration was correlated with successful reinnervation of NMJs, LG muscles from repaired animals were collected and examined for reinner-
KCC2 recovery on the membrane of LG motoneurons (Table 4; Fig. 6F).

**KCC2 depletion is independent of microglia.**

It is possible that microglia are responsible for the induction of KCC2 loss, while connection with the muscle is required for KCC2 restoration. After PNI, BDNF is up-regulated in microglia (Trang et al., 2009), and a large number of studies have reported on the regulation of spinal KCC2 after nerve injuries by microglia-derived BDNF (Coull et al., 2005; Trang et al., 2009; Zhou et al., 2011, 2014; Zhang et al., 2012; Ferrini and De Koninck, 2013; Gomes et al., 2013). To examine whether microglia-BDNF is involved, we used Cx3cr1CreERT/+::bdnf<sup>fl/fl</sup> animals in which bdnf is removed specifically from microglia by cre recombination after tamoxifen treatment. In Cx3cr1CreERT/+ animals, one Cx3cr1 allele is inactivated; therefore, we also included data shown above from WT animals for comparison. All animals (n = 5 WT, 6 Cx3cr1CreERT/+) were analyzed 14 d postinjury and compared to sham operated animals of the same genotype. There were significant differences according only to injury, and not genotype or their interaction (Table 5).

Deletion of BDNF from microglia or heterozygous inactivation of the Cx3cr1 gene have no effect on the down-regulation of KCC2 after axotomy (Fig. 7A).

To further examine a possible role of microglia independent of BDNF, we attenuated the microglial reaction in the ventral horn by removing the CSF1 gene (csf1<sup>f</sup>) from motoneurons using csf1 floxed alleles and a Chat-iRES-Cre mouse line expressing cre in ChAT-expressing neurons (including all motoneurons). This manipulation was previously shown to prevent ventral horn microgliosis while preserving dorsal horn microglial activation after sciatic nerve injury (Rotterman et al., 2019), and we replicated this effect (Fig. 7B). As in WT animals, Chat<sup>iREScre/+::csf1<sup>flo</sup> animals and Chat<sup>iREScre/-::csf1<sup>fl</sup></sup> (n = 5 per group) expressed high levels of KCC2-IR on the cell body of motoneurons contralateral to the injury, while motoneurons axotomized after sciatic nerve cut and ligation were similarly depleted in animals of both genotypes (Table 5).

Thus, attenuating the microglial reaction in the ventral horn after nerve injury did not alter motoneuron KCC2 down-regulation after axotomy (Fig. 7C). We used animals of both sexes (n = 2 females, n = 3 males per group), and both similarly lose KCC2 with or without ventral horn microglia activation. In conclusion, mechanisms that deplete KCC2 on spinal cord motoneurons following axotomy are independent of microglia in both sexes. This is different from mechanisms inducing KCC2 depletion in dorsal horn neurons following nerve injuries, which involve sex-dependent microglia activation. (Maplebeck et al., 2017).

**KCC2 depletion is independent of motoneuron-BDNF or TrkB activation**

Like microglia, axotomized motoneurons also increase BDNF expression after nerve injuries (Kobayashi et al., 1996; Al-Majed et al., 2000a, 2004). To determine whether autocrine BDNF signaling may induce KCC2 downregulation in axotomized motoneurons, we used Chat<sup>iREScre/+::bdnf<sup>fl/fl</sup></sup> mice to knock out bdnf from all cholinergic neurons.

### Table 4

| reps | Reps | SEM | SEM | SEM |
|-------|------|-----|-----|-----|
| 5     | 5    | 98.2| 98.2| 98.2|

### Table 5

| reps | Reinnervation % | SEM | SEM | SEM |
|-------|-----------------|-----|-----|-----|
| 5     | 50              | 58  | 58  | 58  |

### Figure 6

The disappearance of KCC2 coincides with microglial onset but KCC2 restoration depends on muscle reinnervation. CX3CR1<sup>EGFP/+</sup> animals underwent unilateral sciatic nerve injury and were allowed to survive for various time points. A, Representative images of microglial (green) and KCC2 (white) IR 14 d after sciatic nerve cut/ligation, at the peak of the microglial response. Scale bars = 200 μm. Regions of KCC2 depletion overlap with the area of microglial reactivity. B, Time course of KCC2 downregulation and recovery in CX3CR1<sup>EGFP/+</sup> animals. KCC2 protein levels begin decreasing by 3 d and are significantly reduced by two weeks following injury but are restored to control levels by 60 d in animals in which regeneration was allowed (repaid: Table 4; *p < 0.05, * *p < 0.01*). Error bars = SEM. C, D, Microglial reaction (green) around injured LG motoneurons (blue) 14 d (C) or 60 d (D) after sciatic nerve injury in cut/ligated animals. KCC2-IR loss appears to coincide with the onset of microgliosis but persists after the response has attenuated. E, Representative motor endplates in the LG muscle following sciatic nerve transection and repair. Acetylcholine receptors were labeled with α-bungarotoxin (red) and motor axons were identified by VAChT (white) and NF-H (green). Scale bars = 5 μm. Endplates with overlapping acetylcholine receptors and VAChT were designated as reinnervated. F, Correlation between KCC2-IR and endplate reinnervation of individual animals (circles). Animals with higher levels of reinnervation of the LG have higher levels of KCC2 fluorescence on LG motoneurons (linear regression; Table 4).
Table 5. Details of statistics for experiments manipulating microglia and motoneuron BDNF production

| Data reference | Data structure | Type of test | Power and statistical significance |
|----------------|---------------|--------------|-----------------------------------|
| **A**<br> WT (intact) vs WT (cut/ligated) | Normal distribution | Two-way ANOVA | Power = 1.0 |
| ... | ... | ... | ... |
| **B**<br> Chat<sup>RESCre</sup>−/−:csf1<sup>fl/fl</sup> (intact) vs Chat<sup>RESCre</sup>−/−:csf1<sup>fl/fl</sup> (cut/ligated) | Normal distribution | Two-way ANOVA | Power = 0.998 |
| ... | ... | ... | ... |
| **C**<br> Chat<sup>RESCre</sup>−/−:bdnf<sup>+/−</sup> (intact) vs Chat<sup>RESCre</sup>−/−:bdnf<sup>+/−</sup> (cut/ligated) | Normal distribution | Two-way ANOVA | Power = 0.994 |
| ... | ... | ... | ... |
| **D**<br> Chat<sup>RESCre</sup>−/−:bdnf<sup>+/−</sup> (intact) vs Chat<sup>RESCre</sup>−/−:bdnf<sup>+/−</sup> (cut/ligated) | Normal distribution | Two-way ANOVA | Power = 0.05 |
| ... | ... | ... | ... |

Boldface indicates the group against all other comparisons are made in multiple comparisons tests.

Removal of BDNF from motoneurons did not impact baseline KCC2 levels or KCC2 decrease after axotomy (Fig. 8A). We compared Chat<sup>RES-Cre</sup>−/−:bdnf<sup>/−</sup> (n = 4) and Chat<sup>RES-Cre</sup>−/−:bdnf<sup>+/−</sup> animals (n = 4) and found no significant difference in KCC2-IR levels between uninjured motoneurons and similar statistically significant reductions in injured motoneurons in animals of both genotypes (Table 5).

In these experiments, all cholinergic neurons lacked BDNF throughout development, raising the possibility of compensatory mechanisms. To circumvent this potential pitfall, we analyzed a cohort of animals using SLICK mice. We used specifically the SLICK-A line that we have used previously to delete bdnf in a cell-specific manner (Krawkowiski et al., 2015). In these animals, YFP and CreER<sup>T2</sup> are constitutively expressed under the control of the thy1...

Figure 7. KCC2 depletion occurs independent of microgliosis. A, Relative KCC2 depletion in WT and CX3CR1 BDNF KO animals 14 d after cut/ligation. Genotype had no effect on KCC2 levels in motoneurons contralateral or ipsilateral to injury (Table 5). Removing BDNF from microglia had no impact on KCC2 expression. B, Lumbar spinal cord sections from animals expressing normal CSF1 (top) and with CSF1 removed from motoneurons (bottom) 14 d after ligation. Chat<sup>RESCre</sup>−/−:csf1<sup>fl/fl</sup> animals (bottom image) exhibit normal microgliosis in the dorsal horn but have the microglial reaction to injury greatly attenuated in the ventral horn compared to Chat<sup>RESCre</sup>−/−:csf1<sup>fl/fl</sup> animals (top image). Scale bars = 200 μm. C, KCC2 immunofluorescence 14 d after sciatic ligation in csf1<sup>fl/fl</sup> animals. Preventing microgliosis in the ventral horn had no impact on KCC2 within intact or injured motoneurons (Table 5). Error bars = SEM; **p ≤ 0.01, ***p ≤ 0.001.
cells are axotomized LG motoneurons, but only YFP+ and FB+ motoneurons would have undergone cre recombination removing BDNF expression. Due to the sparse YFP/cre expression of motoneurons, 10 YFP/cre+ neurons per animal were not feasible even after doing bilateral injections and axotomy to increase the number of FB/ YFP/cre-expressing motoneurons. Membrane KCC2-IR was measured in 2–10 FB/YFP+ motoneurons per animal (4.23 ± 2.75 average neurons/animal ± SD) to obtain each animal average and compared to similar numbers of FB/non-YFP (cre–) neurons (n = 2–10 neurons, average 4.22 ± 3.35). We compared SLICK: : bdnf+/− animals (n = 3 animals) with SLICK: : bdnf−/− animals (n = 5 animals) all treated with tamoxifen before FB retrograde labeling and bilateral sciatic nerve cut and ligation. There was no significant difference in KCC2 levels between injured FB+/YFP+ and FB+/YFP− motoneurons (with or without cre, respectively) in either genotype 14 d after injury (Fig. 8B,C; Table 5). Removing BDNF from adult motoneurons did not prevent KCC2 downregulation, similar to the lack of effects of removing BDNF from all motoneurons throughout development.

**KCC2 depletion is independent of TrkB activation**

Removing BDNF from either of the two main sources in the spinal cord after injury did not have any effects on KCC2 regulation, but it is still possible that another source could compensate for motoneuron or microglial BDNF. Moreover, removal of BDNF could also be compensated for by upregulation of neurotrophin-4/5, another ligand of TrkB. It has also been shown that ligand-independent autophosphorylation of TrkB receptors can occur (Zaccaro et al., 2001). We therefore blocked the TrkB receptor directly with either the competitive TrkB antagonist ANA-12 (Cazorla et al., 2011) or by using the F616A mutant pharmacogenetic system (Chen et al., 2005).

In ANA-12 experiments, mini-osmotic pumps were used to continuously deliver the drug or vehicle intraperitoneally into WT animals, starting at the time of injury and until euthanasia (14 d). Systemic ANA-12 delivered via intraperitoneal injections has been shown to cross the blood brain barrier and effectively inhibit TrkB 4 h post-injection (Cazorla et al., 2011). The concentration of ANA-12 in the pump was adjusted such that we delivered 1 ng/h into the animal. ANA-12 actions were confirmed with Western blotting by probing for pERK, a target of TrkB. It has also been shown that ligand-independent autophosphorylation of TrkB receptors can occur (Zaccaro et al., 2001). We therefore blocked the TrkB receptor directly with either the competitive TrkB antagonist ANA-12 (Cazorla et al., 2011) or by using the F616A mutant pharmacogenetic system (Chen et al., 2005).

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Promoter in sparse populations of neurons, including some motoneurons. CreER<sup>T2</sup> is activated in the presence of tamoxifen, which we administered in adulthood, thereby eliminating developmental confounds. This approach has the added advantage of allowing internal controls since only YFP+ cells have the capacity of cre recombination, leaving YFP− populations in the same animals unaffected. We crossed these animals with bdnf<sup>−/−</sup> animals and retrogradely labeled LG motoneurons with FB. Thus, all FB+...
underwent a sham injury in one side. In sham control animals, no significant differences in KCC2 membrane immunofluorescence were found on the sham side compared to the contralateral side, and the levels of KCC2-IR was comparable to all control motoneurons in previous experiments. This suggests that basal expression of KCC2 was not affected by ANA-12 treatment. In injured animals, we found similar reductions of KCC2 in injured motoneurons in animals treated with either vehicle (n = 4) or with ANA-12 (n = 6) compared to control motoneurons (Fig. 9B). There were no significant differences in sham injured and intact motoneurons or reductions in KCC2 levels between injured neurons treated with ANA-12 or vehicle (Table 6). In conclusion, ANA-12 treatment did not prevent KCC2 downregulation in axotomized motoneurons.

ANA-12 is a partial competitive antagonist of TrkB (Chen et al., 2015) and therefore we cannot assume block of ligand binding to be complete. To obtain further proof of a lack of a role for TrkB in KCC2 downregulation, we blocked TrkB kinase activity (and thus signaling) using F616A mutant mice. F616A animals express a TrkB mutation with a single base substitution that makes it sensitive to the kinase inhibitor 1NMPP1 (Chen et al., 2005). 1NMPP1 was administered in the drinking water using established protocols that prevent TrkB downstream signaling (Chen et al., 2005; Wang et al., 2009; Rantamäki et al., 2011). In agreement with ANA-12 experiments, in F616A animals treated with normal drinking water (n = 3) and 1NMPP1 (n = 6) normal levels of KCC2 were found on uninjured FB LG motoneurons. A downregulation of KCC2 was also found after injury in FB LG motoneurons regardless of treatment (Table 6; Fig. 9C). Taken together, the results in ANA-12-treated animals and F616A animals strengthened the conclusion that TrkB signaling is not involved in KCC2 regulation after sciatic nerve injury.

Table 6. Details of statistics for experiments blocking TrkB signaling

| A | Data reference | Data structure | Type of test | Power and statistical significance |
|---|----------------|----------------|--------------|-----------------------------------|
| 7B | Intact ANA-12 vs sham ANA-12 | Normal distribution, unequal variance | Two-way ANOVA | Power = 0.608 |
|   | Injured VEH | | Side; $F_{1,27} = 6.474; p = 0.018$ | Power = 0.993 |
|   | Intact ANA-12 | | Treatment; $F_{2,27} = 13.095; p < 0.001$ | Power = 0.860 |
|   | Injured ANA-12 | | Interaction; $F_{2,27} = 7.289; p = 0.004$ | $p = 0.576$ |
|   | Post hoc Bonferroni Injured VEH | | | $p > 0.999$ |
|   | Post hoc Bonferroni Intact ANA-12 | | | $p = 0.003$ |
|   | Post hoc Bonferroni Injured ANA-12 | | | $p > 0.999$ |
|   | Post hoc Bonferroni | | | $p = 0.006$ |
| B | 7C | Normal distribution | Two-way ANOVA | Power = 1.00 |
|   | Intact VEH vs injuried VEH | | Side; $F_{1,17} = 35.783; p < 0.001$ | Power = 0.05 |
|   | Intact 1NMPP1 | | Treatment; $F_{1,17} = 0.935; p = 0.350$ | Power = 0.079 |
|   | Injured 1NMPP1 | | Interaction; $F_{1,17} = 1.325; p = 0.269$ | $p = 0.002$ |
|   | Post hoc Bonferroni Injured VEH | | | $p = 0.470$ |
|   | Post hoc Bonferroni Intact ANA-12 | | | $p < 0.001$ |

Boldface indicates the group against all other comparisons are made in multiple comparisons tests.
allow us to conclude that KCC2 downregulation in axotomized motoneurons is independent of TrkB signaling.

**Discussion**

In this study, we demonstrate that KCC2-IR is significantly decreased on the cell membranes of spinal motoneuron somata and dendritic processes following peripheral axotomy. KCC2 disappearance begins with a downregulation of KCC2 mRNA in the days immediately after injury, followed by a slower disappearance of protein from the membrane lasting weeks. We also found that KCC2 is restored only after successful axon regeneration and muscle reinnervation, raising the possibility that peripheral, rather than central, signals may regulate KCC2 expression. Furthermore, KCC2 regulation on peripherally axotomized motoneurons occurs independent of microglia and BDNF/TrkB signaling. This is different from the canonical mechanism for KCC2 downregulation that has been established in dorsal horn neurons. Instead, KCC2 downregulation on spinal motoneurons is more similar in time course and extent to what has been previously reported in brainstem motor nuclei following peripheral axotomy.

**KCC2 is preferentially preserved in distal dendrites**

Retention of KCC2 in the distal dendrites of spinal motoneurons, despite strong reductions in KCC2 mRNA, suggests differences in KCC2 stability in different dendrite compartments. Dendrites also lose most synapses on proximal, rather than distal, compartments after axotomy (Brännström and Kellerth, 1998; Rotterman et al., 2014). Interestingly, electrophysiological studies have also shown that glycinegic (strychnine-sensitive) synapses, found mostly on the somata (Takata and Ogata, 1980) sustained larger decreases in efficacy than more distally located GABAergic (picrotoxin-sensitive) synapses (Takata and Ogata, 1980; Takata and Nagahama, 1983). It is worth noting that in some central neurons, distal dendrites are enriched with the KCC2a isoform (Markkanen et al., 2014) and that KCC2 in dendrites is frequently associated with the maintenance of structural properties at excitatory synapses and less with a chloride transport function (Gulyás et al., 2001; Blaesse and Schmidt, 2015). Recent data from using isotype specific antibodies (P. Calvo and F. J. Alvarez, unpublished observations) shows that KCC2b is uniquely expressed on the somatic and proximal dendrite membrane of spinal motoneurons. Possible differences in KCC2 isoforms, as well as anchoring, membrane stabilization, and function in distinct dendritic compartments are all possibilities for explaining the differential loss of KCC2 in motoneuron soma and dendrites that deserve further study.

It should also be noted that, for the first time, we have reported trafficking of kcc2 mRNA into dendritic compartments of motoneurons. KCC2 transcription of kcc2 mRNA within the dendrites is one alternative explanation for the differential regulation of dendritic KCC2. Our Nissl-stained sections only permitted analyses of proximal dendrites as they emerge from the cell body. Future studies should be performed combining RNA-Scope kcc2 mRNA detection and dendritic labeling with AAV1 transfection to localize kcc2 mRNA throughout the dendritic arbor. The high sensitivity and specificity of the RNA-Scope technique would allow detection of small quantities of mRNAs in dendritic compartments. Nonetheless, we believe it is unlikely that there is enough preservation of kcc2 mRNAs in distal dendrites to maintain KCC2 protein; the scattered dots of mRNA signal in the neuropil adjacent to the axotomized motoneurons also disappeared after injury. Therefore, the best explanation at present is that KCC2 in the very distal dendrites has a very low turnover rate, explaining their preservation at the time point of our analyses. In this case, we would expect chronic axotomy at longer time points would result in the eventual loss of this KCC2 as well.

It is also possible KCC2 in distal dendrites is serving a different purpose. Independent of its role in ion transport, KCC2 has been shown to stabilize excitatory synapses and be important in dendritic spine development by regulating actin phosphorylation and stabilization (Li et al., 2007; Liano et al., 2015). KCC2-deficient mice have fewer functional glutamatergic synapses and excessively long dendritic protrusions that cause an increase in the total dendrite length of pyramidal cells (Li et al., 2007). The regions where we see KCC2 preservation on motoneurons (the last 75–100 µm of dendrite) typically have very low synaptic densities (Alvarez et al., 1997), making synapse stabilization an unlikely explanation for KCC2 retention there. However, these most distal regions of the dendrites undergo many changes after long-term axotomy without regeneration, including becoming more highly branched, increasing in diameter, and becoming axon-like in both structure and molecular composition (Rose and Odlozinski, 1998; Rose et al., 2001; MacDermid et al., 2002). At the time points we observed, no drastic changes in distal dendrite morphology were evident, but it is possible that the high density of KCC2, if lost long-term, is contributing to the eventual morphologic changes observed with longer-term axotomy (Rose and Odlozinski, 1998).

**KCC2 regulation in axotomized motoneurons**

The first study of KCC2 regulation on motoneurons following PNI was conducted in the dorsal motor nucleus of the vagus (DMV), where KCC2 mRNA was diminished by 3 d after vagal nerve axotomy in young rats (Nabekura et al., 2002). DMV neurons innervate postganglionic parasympathetic neurons rather than skeletal muscle, but these findings were later recapitulated in facial (Toyoda et al., 2003; Kim et al., 2018) and hypoglossal (Tatetsu et al., 2012) motoneurons following transaction of the respective peripheral nerves. Several of these studies speculated on the eventual return of KCC2 mRNA (Toyoda et al., 2003) and protein (Tatetsu et al., 2012; Kim et al., 2018), but none analyzed this directly or whether it was dependent on successful target reinnervation. Our study established that KCC2 recovery in axotomized motoneurons is correlated with successful axon regeneration and more specifically with re-innervation of NMJs, this mechanism could be shared by both brainstem and spinal motoneurons.
While brainstem and spinal motoneurons both innervate muscle, it should be noted that the circuitry surrounding them is vastly different and responses to injury may not always be equivalent across motoneuron populations. Whereas facial (Vaughan, 1994; Kim et al., 2018), like spinal motoneurons (Alvarez et al., 2011; Rotterman et al., 2019) have small decreases in gephyrin+ (GABA/glycinergic) synapses following peripheral axotomy, hypoglossal motoneurons do not lose inhibitory synapses (Sumner, 1975; Tatetsu et al., 2012). Interestingly, inhibitory synapses on facial motoneurons increased in size and GABA content following axotomy (Vaughan, 1994). While this has not been investigated on spinal motoneurons following PNI, it is possible that they also shift their inhibitory synapse phenotype or strength following injury. Many inhibitory synapses in the spinal cord are mixed GABA/glycine synapses, and they shift from predominantly GABAergic to predominantly glycinergic during postnatal development (Gao et al., 2001; González-Forero and Alvarez, 2005). Thus, adult spinal motoneurons predominantly receive inhibition from glycinergic synapses (Alvarez et al., 1997) but could revert to GABAergic mechanisms after axotomy. Postsynaptic GABA_A synaptic currents have much slower decay times than glycinergic synapses (Alvarez, 2017), allowing for longer depolarizations in the context of diminished KCC2 expression and high internal chloride. During development, while KCC2 levels are low, high internal chloride and long-lasting GABAergic depolarizations drive neurite outgrowth (Sernagor et al., 2010). Thus, in contrast to dorsal horn neurons after PNI or motoneurons following SCI (which highlight disinhibition after KCC2 loss), KCC2 changes on axotomized brainstem motoneurons seems to induce a state in which GABAergic transmission is the direct driver of activity. Correspondingly, loss of KCC2 was coupled with increased calcium levels and firing in response to GABA exposure in slice preparations of axotomized DMV motoneurons (Nabekura et al., 2002) and with spontaneous calcium oscillations in axotomized facial motoneurons that were driven by depolarizations dependent on activation of GABA_A and NMDA receptors (Toyoda et al., 2003).

KCC2 regulation after PNI differs between motoneurons and dorsal horn interneurons

Like motoneurons, dorsal horn spinal neurons (more specifically nociceptive Lamina I neurons) decrease KCC2 following nerve injuries, resulting in disinhibition and behavioral hyperalgesia (Coull et al., 2003, 2005). Unlike motoneurons, dorsal horn neurons are fully contained within the CNS and are not directly injured or axotomized after PNI. Instead, signals from injured sensory afferents trigger the activation of microglia in the dorsal horn and subsequent release of BDNF that depletes KCC2 in neighboring neurons, at least in males (Tsuda et al., 2003; Coull et al., 2005; Trang et al., 2009, 2012; Beggs et al., 2012; Sorge et al., 2015). Dorsal horn neurons in female mice also decrease KCC2 but through a different mechanism than males that is not yet well elucidated (Maplebeck et al., 2018). Independent of sex, the changes in KCC2 occur quickly following injury, most typically a constriction of the nerve, and are likely due to changes in KCC2 phosphorylation and membrane trafficking. In axotomized motoneurons, however, KCC2 is downregulated at the transcriptional level and it takes two to three weeks for complete loss of the protein from the membrane, likely because of slow protein turnover.

Another major distinction between spinal motoneurons after PNI and mechanisms described in other models is a differential dependence on microglia and BDNF. BDNF-TrkB can regulate KCC2 membrane content by post-translational mechanisms, altering KCC2 phosphorylation, or directly by regulating KCC2 gene expression (for review, see Lee-Hotta et al., 2019). The cell bodies of axotomized motoneurons are quickly surrounded by a layer of activated microglia (Aldskogius, 2011) that, similar to dorsal horn microglia, could be involved in regulating KCC2 expression through BDNF release or other mechanisms. However, our results negate this hypothesis, demonstrating a lack of involvement of ventral microglia on the downregulation of KCC2 in axotomized motoneurons. Microglia activation phenotypes in the dorsal and ventral horn after PNI are not necessarily the same, and differences might further increase depending on the type of injury. Although activation of microglia in the dorsal horn is also dependent on CSF1, this time released from affected primary afferents (Guan et al., 2016), the release of BDNF from activated dorsal horn microglia was shown to specifically depend on microglia P2X4 receptors activated by ATP released from the central terminals of overactive sensory afferents (Beggs et al., 2012). Ventral horn microglia also upregulate several classes of purinergic receptors after nerve injury (Kobayashi et al., 2012), but their exact roles have not yet been investigated. Although ventral horn microgliosis is also dependent on CSF1 released by axotomized motoneurons (Rotterman et al., 2019), some features of the microglia reaction to nerve injury differ between the dorsal and ventral horn (Rotterman et al., 2019). Another significant difference is the type of nerve injury leading to activation of microglia. Dorsal horn studies are focused on a model of partial ligation of the sciatic nerve known to cause a well-characterized hyperalgesia (Bennett and Xie, 1988), while we studied the consequences of full transection of the nerve. One study comparing the phenotype of dorsal horn microglia in these two types of injuries found many significant differences (Hu et al., 2007). In conclusion, the response of microglia in the ventral horn is not involved in the downregulation of KCC2 from motoneurons after nerve transections and is thus different from the well-established role of dorsal horn microglia, at least in males, after hyperalgesia-inducing nerve injuries. 

KCC2 downregulation is independent of TrkB activation

BDNF is transiently upregulated in motoneurons following axotomy (Kobayashi et al., 1996; Al-Majed et al., 2000a, 2004), and the release of BDNF is key in treatments that enhance motor axon regeneration and functional recovery following injury (Al-Majed et al., 2000a,b; English et al., 2004; Coull et al., 2003, 2005). Instead, factors released from activated microglia likely regulate KCC2 in dorsal horn spinal neurons following axotomy.
Comparison of KCC2 downregulation in motoneurons after nerve injury or SCI

Following SCI, motoneurons below the level of injury also partially lose KCC2 (Boulenguez et al., 2010). These motoneurons are not directly injured during SCI and their axons remain intact in the peripheral nerve. A post-translational mechanism has been proposed to explain this effect, indicating that KCC2 downregulation was dependent on the loss of descending serotonergic inputs and 5HT2A receptor activity (Bos et al., 2013). The downregulation of KCC2, as measured in just the cell body, was estimated to be around 20% (but may be better preserved on the dendrites). This lead to a small but significant increase in the reversal potential of chloride of only 5 mV in adult rats (Murray et al., 2011) that was sufficient to generate significant disinhibition and spasticity. After SCI, motoneuron excitability and spasticity can be rectified with enhancers of KCC2 activity (Sánchez-Brualá et al., 2018), suggesting that enough KCC2 remains on the surface of the motoneurons to rescue KCC2 function and the animal from spasticity. In contrast, after axotomy of adult facial motoneurons, $E_{\text{GABA}}$ depolarizes by almost 19 mV (Toyoda et al., 2003). In facial motoneurons and spinal motoneurons, almost no KCC2 mRNA or protein is present two weeks after injury, and we have shown that the loss includes substantial regions of the dendritic arbor. Thus, while downregulation of KCC2 in motoneurons after SCI leads to a partially disinnhibited state, the more complete downregulation we describe here might lead to GABA/glycine synapses reversals climbing above action potential thresholds and becoming significant drivers of activity in regenerating motoneurons. Given the drastic change in KCC2 on axotomized motoneurons and the close link of this loss with regenerative state, it is possible that KCC2 reductions may be tied to motoneuron regenerative mechanisms. Activity-dependent treatments are known to enhance regeneration (Al-Majed et al., 2000a,b; English et al., 2007, 2009; Udina et al., 2011) and this enhancement could be complemented by the many changes in the physiology of regenerating motoneurons that increase their excitability: increased input resistance, decreased rheobase, and changes in action potential dynamics (Kuno et al., 1974; Huizarr et al., 1977; Gustafsson, 1979; Gustafsson and Pinter, 1984; Gonzalez-Forero et al., 2007). In this context KCC2 depletion resulting in enhanced "excitatory" drive from GABA/glycine synapses may further promote motoneuron firing and calcium entry, contributing to regeneration. This is a hypothesis that will need to be fully investigated in the future. It is however important to consider at present the possibility that while KCC2 depletion in the adult spinal cord can be maladaptive in some conditions (SCI spasticity, hyperalgesia), it might be beneficial in other situations (motoneuron regeneration after PNI).

In summary, KCC2 downregulation in spinal interneurons versus motoneurons after PNI and in motoneurons in SCI versus PNI differ in mechanism, completeness, and functional role. This should be taken into account when assessing the potential impact of KCC2-enhancing manipulations after different kinds of injuries.

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