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Overexpression of the Astrocyte Glutamate Transporter GLT1 Exacerbates Phrenic Motor Neuron Degeneration, Diaphragm Compromise, and Forelimb Motor Dysfunction following Cervical Contusion Spinal Cord Injury

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A major portion of spinal cord injury (SCI) cases affect midcervical levels, the location of the phrenic motor neuron (PhMN) pool that innervates the diaphragm. While initial trauma is uncontrollable, a valuable opportunity exists in the hours to days following SCI for preventing PhMN loss and consequent respiratory dysfunction that occurs during secondary degeneration. One of the primary causes of secondary injury is excitotoxic cell death due to dysregulation of extracellular glutamate homeostasis. GLT1, mainly expressed by astrocytes, is responsible for the vast majority of functional uptake of extracellular glutamate in the CNS, particularly in spinal cord. We found that, in bacterial artificial chromosome-GLT1-enhanced green fluorescent protein reporter mice following unilateral midcervical (C4) contusion SCI, numbers of GLT1-expressing astrocytes in ventral horn and total intraspinal GLT1 protein expression were reduced soon after injury and the decrease persisted for ≥6 weeks. We used intraspinal delivery of adeno-associated virus type 8 (AAV8)-Gfa2 vector to rat cervical spinal cord ventral horn for targeting focal astrocyte GLT1 overexpression in areas of PhMN loss. Intraspinal delivery of AAV8-Gfa2-GLT1 resulted in transduction primarily of GFAP - astrocytes that persisted for ≥6 weeks postinjury, as well as increased intraspinal GLT1 protein expression. Surprisingly, we found that astrocyte-targeted GLT1 overexpression increased lesion size, PhMN loss, phrenic nerve axonal degeneration, and diaphragm neuromuscular junction denervation, and resulted in reduced functional diaphragm innervation as assessed by phrenic nerve–diaphragm compound muscle action potential recordings. These results demonstrate that GLT1 overexpression via intraspinal AAV-Gfa2-GLT1 delivery exacerbates neuronal damage and increases respiratory impairment following cervical SCI.

Key words: adeno-associated virus; gene therapy; glutamate transporter; phrenic motor neuron; respiratory; spinal cord injury

Introduction

Cervical spinal cord trauma represents greater than half of all human spinal cord injury (SCI) cases, in addition to often result-

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ondary cell death and consequent exacerbation of functional deficits (Oyinbo, 2011). Glutamate-mediated excitotoxicity plays a key role in these secondary events. Large increases in extracellular glutamate occur shortly after SCI, and elevation can also persist, depending on injury severity (Panter et al., 1990; Liu et al., 1991; McAdoo et al., 1999). Early gray matter loss is likely mediated by NMDA receptors, while delayed loss of neurons and oligodendrocytes, as well as axonal and myelin injury, is thought to be predominantly mediated via AMPA overactivation. Spinal motor neurons are particularly vulnerable to glutamate excitotoxicity due to such factors as expression of calcium-permeable glutamate receptors and low calcium-buffering capacities (Xu et al., 2004, 2008).

Glutamate transporter-1 (GLT1) plays a central role in regulating extracellular glutamate levels. GLT1 is almost exclusively expressed by astrocytes and accounts for ~90% of functional uptake of extracellular glutamate (Tanaka et al., 1997). Following SCI, astrocyte death, and/or compromised GLT1 expression, membrane localization and/or function can result in dysregulation of glutamate homeostasis. Several studies have shown that GLT1 expression is chronically reduced following thoracic SCI (Kim et al., 2011; Min et al., 2012). Changes in GLT1 expression appear to be spatiotemporally regulated in a more subtle manner (Xu et al., 2008; Gwak et al., 2012). Rapid GLT1 upregulation has been reported within the first hours post-SCI (Vera-Portocarrero et al., 2002) in surviving astrocytes (Xu et al., 2008). This increase may function to counteract immediate elevation of glutamate during the early phase; however, this initial response is followed by long-lasting downregulation due to astrocyte death and reduced expression in reactive astrocytes and even in newly generated cells arising from astrocyte proliferation (Xu et al., 2008; Min et al., 2012). These data suggest that the long-term ability of astrocytes to regulate extracellular glutamate homeostasis post-SCI is compromised, which increases susceptibility to secondary excitotoxic damage.

Selective GLT1 overexpression in adult astrocytes had previously made it difficult to clearly determine its exact contribution to neuroprotection following SCI. To this end, we used recombinant adeno-associated viral type 8 (AAV8) vectors to selectively drive overexpression of GLT1 in astrocytes in a clinically relevant model of midcervical contusion SCI. Furthermore, we evaluated key histological and functional outcomes associated with a large proportion of human SCI, namely PhMN loss, diaphragm denervation, and consequent respiratory dysfunction.

Materials and Methods

Animals

Transgenic bacterial artificial chromosome (BAC)-GLT1-enhanced green fluorescent protein (eGFP) transgenic mice were previously generated by Regan et al. (2007) and were a kind gift from Jeffrey Rothstein’s laboratory at Johns Hopkins University. BAC-GLT1-eGFP mice were used to quantify GLT1 promoter activity following cervical contusion. Male BAC-GLT1-eGFP mice and wild-type C57BL/6 mice aged from 8 to 12 weeks were used in the study. Female Sprague Dawley rats weighing 250–300 g were purchased from Taconic Farm. Rats were divided into three paradigm groups according to the experimental design described below. All animals were housed in a humidity-controlled, temperature-controlled, and light-controlled animal facility under standard chow and with ad libitum access to water. Experimental procedures were approved by the Thomas Jefferson University institutional animal care and use committee and conducted in compliance with the European Communities Council Directive (2010/63/EU, 86/609/EEC, and 87-848/EEC), the National Institutes of Health Guide for the Care and Use of Laboratory Animals, and the Society for Neuroscience’s Policies on the Use of Animals in Neuroscience Research.

Experimental design

Mouse SCI paradigm. BAC-GLT1-eGFP mice were subjected to cervical contusion SCI at both C4 and C5 levels (Nicaise et al., 2012b), and they were then killed 2, 14, or 42 d postinjury (DP1). Spinal cord tissue was processed for histology to quantify the number of eGFP+ cells in the ventral horn ipsilateral to the lesion site. Nontransgenic C57BL/6 mice also received C4 C5 spinal contusion to measure whole-tissue GLT1 protein levels by immunoblotting. After the animal was killed, the spinal cord was freshly removed, subdivided into 1.5-mm-long hemicord fragments encompassing the epicenter of the injury or spinal cord segments caudal to the epicenter (Fig. 1C), and then snap-frozen with liquid nitrogen.

Rat SCI paradigm 1. AAV-Gfa2-eGFP or AAV-Gfa2-GLT1 was injected into the spinal cord ventral horn of sham (laminectomy-only uninjured) or spinally contused rats, and rats were then killed 2 or 10 d postinjection. This preliminary study was used to determine the efficiency of the virus for targeting/transducing spinal cord astrocytes, the timing of GLT1 overexpression, and the spatial distribution/spread of virus transduction.

Rat SCI paradigm 2. AAV-Gfa2-eGFP or AAV-Gfa2-GLT1 was injected into the spinal cord ventral horn at the C3, C4, and C5 levels ipsilateral to the injury site (see Fig. 3A) 10 d before unilateral C4 contusion (see Fig. 5A). Rats were killed 42 DPI. This “pretreatment” paradigm was aimed at boosting the levels of GLT1 within the spinal cord before injury as proof of principle. These animals were used for comprehensive behavioral, electrophysiological, and histological analysis of the therapeutically efficacious GLT1 overexpression.

Rat SCI paradigm 3. Rats were injured at the C4 level and were then immediately injected with AAV-Gfa2-eGFP or AAV-Gfa2-GLT1 into spinal cord ventral horn at the C3, C4, and C5 levels ipsilateral to the injury site (see Figs. 3A, 8A). This “acute” AAV injection was tested as a more clinically relevant paradigm of AAV-mediated GLT1 overexpression.

Cervical contusion SCI

Cervical contusion SCI was performed on mice and rats. Mice were anesthetized with ketamine (120 mg/kg) and xylazine (5 mg/kg). The dorsal skin and underlying muscle layers were incised and retracted between the spinous processes of C2 and T1. The paravertebral muscles overlying C4–C6 were removed. Following right-sided unilateral laminectomy at C4 and C5 levels, mice were subjected to a double spinal contusion injury with the Infinite Horizon impactor (Precision Systems and Instrumentation) at both C4 and C5 levels using a 1 mm tip. The force of both impacts was 50 kdyn (impact on C4 was generated first). Sham control mice underwent the same surgery, including laminectomy, but did not receive contusion injury. Rats were anesthetized with ketamine (100 mg/kg), xylazine (5 mg/kg), and acepromazine (2 mg/kg). The cervical dorsal skin and underlying muscles were incised. The paraspinal muscles overlying C4 were retracted. An 1.5 × 10⁶ cells were seeded into 150-mm Petri dishes in DMEM supplemented with 10% FBS, 1 mM MEM sodium pyruvate, 0.1 mM MEM nonessential amino acids, and 0.05% penicillin-streptomycin (5000 U/ml). At 24 h
after cell seeding, medium was changed to culture medium containing only 5% FBS, and cells were transfected with three plasmids: adenovirus helper plasmid (pEΔ6), AAV helper encoding serotype 8 (kindly provided by Miguel Sene Esteves at the University of Massachusetts), and the AAV packaging vector containing the glycolytic acidic protein (GFPa) promoter flanked by either the eGFP gene or the human EAA2 gene sequence (obtained from Jeffrey Rothstein’s laboratory). Plasmids were transfected into HEK293T cells using Polyfect according to the manufacturer’s specifications (Qiagen). Cultures were incubated at 37°C in 95% O2 and 5% CO2. After 72 h, cells were harvested and pelleted by centrifugation. The pellet was resuspended in 10 mM Tris, pH 8.0, and chilled on ice. Cells were lysed by repeated freeze–thaw cycles, followed by treatment with 50 U benzonase (Novagen) and 0.5% sodium deoxycholate for 30 min at 37°C. The lysate was sonicated three times. Virus was then purified by density gradient centrifugation in iodixanol according to the method of Zolotukhin et al. (1999). Each virus preparation was sterile filtered, and the titer (genomic particles/ml) was determined by preparation was sterile filtered, and the titer (genomic particles/ml) was determined by quantitative RT-PCR using primers and a probe specific for the woodchuck posttranscriptional regulatory element sequence.

Virus delivery was conducted on deeply anesthetized rats. Following unilateral right-sided laminectomy, three injections, each containing 1 µl of virus (titer of 1012 genomic particles/ml), were administered in the spinal cord ventral horn at levels C3, C4, and C5 using a Hamilton gas-tight syringe mounted on a World Precision International; Lepore and Maragakis, 2011; Lepore, 2011). As shown in Figure 3A, the sites of injections were targeted along the rostral–caudal axis with a virtual line connecting the points of dorsal root entry (dotted line) and in the lateral axis corresponding to the middle of each spinal segment (C3, C4, or C5). Ventral horns were targeted by lowering the 33 gauge × 0.5 mm bevelled needle 1.5 mm below the dorsal surface of the spinal cord. Each injection was delivered at a constant rate over 5 min. Upon completion of virus delivery, a synthetic biocompatible membrane (BioBrane, UDL Laboratories) was placed on the top of the laminectomy, which allowed for easy access to the spinal cord below the laminectomy during the subsequent spinal contusion surgery (in experimental rat paradigm 2). Overlying muscles were then closed in layers with sterile 4-0 silk sutures, and the skin incision was closed using sterile wound clips. Animals were allowed to recover and monitored daily.

**Retrograde anatomical tracing of phrenic motor neurons**

Two weeks after the rats were killed, cholera toxin beta fragment (CTβ) conjugated to Alexa555 (Life Technologies) was injected unilaterally into the intrapleural space of the ipsilateral hemidiaphragm muscle (Nicaise et al., 2012a). This procedure allowed for the specific labeling of PhMNs innervating the diaphragm through retrograde transport of CTβ fluorescent tracer. Following laparotomy under anesthesia, 15 µl of CTβ (0.2% solution in distilled water) was intrapleurally delivered through the diaphragm (three injections of 5 µl each) using a 20 µl Hamilton syringe with an attached 33 gauge needle. Abdominal muscles were then sutured with 4-0 sutures, and skin was closed with wound clips.

**Tissue processing for histology**

At the time the animals were killed, they were anesthetized, and diaphragm muscle and phrenic nerves were freshly removed before paraformaldehyde perfusion and then further processed for nerve histology and neuromuscular junction (NMJ) labeling. Animals were transcardially perfused with 0.9% saline, followed by 4% paraformaldehyde perfusion and then further processed for nerve histology and neuromuscular junction (NMJ) labeling. Spinal cords were harvested, then cryoprotected in 30% sucrose for 3 d and embedded in freezing medium. Spinal cord tissue blocks were cut serially in the sagittal or transverse planes at a thickness of 30 μm. Sections were collected on glass slides and stored at −20°C until analysis. Spinal cord sections were thawed, allowed to dry for 1 h at room temperature, and stained with 0.5% cresyl violet acetate according to standard procedure (Nicaise et al., 2012a). For CTβ and eGFP cell counting, frozen spinal cord sections were air-dried, washed, and coverslipped with fluorescent-compatible mounting medium (ProLong Gold, Life Technologies).
Phrenic nerve histology

Phrenic nerves were processed as previously described (Kim et al., 2011). Upon removal, phrenic nerves were fixed in 4% glutaraldehyde solution for 1.5 h, thoroughly washed in Millonig’s buffer with 0.5% (w/v) glucose for 24 h, postfixed in 2% (w/v) OsO₄ for 30 min, dehydrated, and embedded in Epoxy resin LX112. Semithin cross sections (1–2 μm thick) were obtained and stained with 1% toluidine blue (Sigma-Aldrich) diluted in 2% sodium borate solution. The total nerve area and the total number of axons and their cross-sectional areas were assessed by two investigators (T.J.H. and C.N.) in a blind manner using ImageJ software, as previously reported (Leroy et al., 2007).

NMJ analysis

Fresh hemidiaphragm muscle was dissected from each animal for whole-mount immunohistochemistry, as described previously (Wright and Son, 2007). Animals that did not receive CTB injections were used for this study to avoid overlap of Alexa555-conjugated CTB with the rhodamine-conjugated α-bungarotoxin used to label the nicotinic acetylcholine receptors. To enable antibody penetration, muscle was stretched, pinned down to Sylgard medium (Fisher Scientific), and extensively cleaned to remove any connective tissue. Motor axons and their terminals were labeled with SMI-312 (DSI; Developmental Studies Hybridoma Bank), respectively, and both primary antibodies were detected with fluorescein isothiocyanate-conjugated (FITC) anti-mouse IgG secondary antibodies (Jackson ImmunoResearch Laboratories). Postsynaptic acetylcholine receptors were labeled with rhodamine-conjugated α-bungarotoxin (Life Technologies). Labeled muscles were analyzed for total numbers of NMJs, fully innervated NMJs, partially denervated NMJs, completely denervated NMJs, NMJs with thin preterminal axons, and multiply-innervated NMJs. The latter two are indicative of motor axon regeneration. Whole-mounted diaphragms were imaged on a Fluoview FV1000 confocal microscope (Olympus).

Immunohistochemistry

Frozen spinal cord sections were air-dried, washed, permeabilized with 0.4% Triton X-100 in PBS for 5 min at room temperature, and then incubated in blocking solution (PBS containing 10% normal goat serum and 0.4% Triton X-100) for 1 h at room temperature. Sections were labeled overnight at 4°C with the primary antibodies in blocking solution. Sections were then washed three times with PBS (5 min per wash) and incubated with secondary antibodies in blocking solution for 1 h at room temperature. After washing twice with PBS (10 min per wash), sections were coverslipped. A number of primary antibodies were used. Mouse anti-GLT1 (1:200; EMD Millipore) was used to label neurons (Jackson ImmunoResearch Laboratories). Rabbit anti-Olig2 (1:200; EMD Millipore) was used to label cells of the oligodendrocyte lineage (Lepore et al., 2008a). Rabbit anti-Newsletter (1:200; EMD Millipore) was used to label neurons (Lepore and Fischer, 2005). Rabbit anti-Gal-His (1:50; Cell Signaling Technology) labeled the His-tag fused to GLT1 protein. Rabbit anti-GLT1 (1:800) and mouse anti-GLT1 (1:200) were used to label GLT1 protein (both were provided by Jeffrey Rothstein’s laboratory; Lepore et al., 2008b). Rabbit anti-Newsletter (1:200; Thermo Fisher Scientific) labeled proliferating cells (Lepore et al., 2008a). Secondary antibodies included FITC goat-anti-mouse IgG, FITC goat-anti-rabbit IgG, tetramethylrhodamine isothiocyanate (TRITC) goat-anti-mouse IgG, TRITC goat-anti-rabbit IgG, Alexa Fluor 647 goat-anti-mouse IgG, Alexa Fluor 488 goat-anti-rabbit IgG. All secondary antibodies were purchased from Jackson ImmunoResearch Laboratories and diluted at 1:200 to recognize the matched primary antibody.

Imaging and quantification

Images were acquired with either a Zeiss Imager M2 upright microscope or a Fluoview FV1000 confocal microscope (Olympus) and then analyzed with ImageJ or MetaMorph software. In SCI paradigms, lesion size was quantified in cresyl violet-stained sections (Nicaise et al., 2012a). Specifically, lesion area was determined in every fifth section by tracing both the total area of the hemispinal cord ipsilateral to the contusion site and the actual lesion area. Lesion was defined as areas including both lost tissue (cystic cavity formation) and surrounding damaged tissue in which the normal anatomical structure of the spinal cord was lost. The lesion epicenter was defined as the section with the largest percentage lesioned tissue (relative to total tissue area in the same section). Lesion volume was determined using the Cavalieri estimator of volume ($V = \frac{1}{2} \sum_{i=1}^{n} D \cdot A_{i} \cdot H$), where $A_{i}$ is the area of the lesion, $D$ is the distance between measurements (150 μm), and $H$ is the thickness (30 μm) of each section). Ventral horn motor neurons were outlined in these same cresyl violet-stained sections, and the number of motor neuron cell bodies was quantified in a blind manner. Large motor neurons with a clearly identifiable nucleus and a cell soma >200 μm² (Lepore et al., 2011b) or medium-sized neurons (150–200 μm²) were counted. CTB + PhMN motor neurons were counted in every fifth section across the entire cervical spinal cord and plotted at specific distances relative to the lesion epicenter (Nicaise et al., 2013).

In BAC-GLT1-eGFP mice, eGFP + cells were identified in the ventral horn using MetaMorph software (Molecular Devices). Cells were counted and plotted at specific distances relative to lesion epicenter (i.e., 1.5 and 3.0 mm from the epicenter in both rostral and caudal directions; Lepore et al., 2011a). To quantify GFAP fluorescence signal in the ventral horn, integrated intensities were measured in MetaMorph. GFAP intensity was calculated to be the GFAP-integrated intensity minus the background-integrated intensity of the ventral horn. All GFAP images were acquired using the same exposure settings. In uninjured sham or injured rats injected with AAV-Gfa2-eGFP, numbers of eGFP + cells and rostral-caudal spreading of virus transduction were similarly analyzed along the length of the cervical spinal cord.

In vivo astroglial scar border formation analysis, following immunostaining with mouse anti-GLT1 antibody, 3–4 images per animal (at locations surrounding the injury epicenter) were obtained at ×5 magnification. GFAP + astroglial processes immediately adjacent to the lesion, oriented parallel to the border of lesion, and interwoven and overlapped extensively were regarded as the mature border of the astroglial scar (Wanner et al., 2013). Quantification results are presented as the percentage of astroglial scar border length compared with the length of total lesion border.

GLT1 immunoblotting

Whole-tissue spinal cord samples (1.5 mm length) were homogenized on ice in 0.5 ml of radioimmunoprecipitation assay buffer containing the following (in mM): 50 Tris-HCl, pH 7.6, 150 NaCl, 2 EDTA, 0.1% SDS, 0.01% NP-40, Protease Inhibitor Cocktail (Roche Diagnostics), Equal amounts of protein (determined by Bradford assay) were resolved by SDS-PAGE on 4–12% gels and transferred to nitrocellulose membranes. The membranes were blocked in Odyssey blocking buffer (Li-Cor) for 1 h at room temperature and probed with anti-GLT1 antibody (1:5000; Li-Cor) as secondary antibody for detection. Acquisition was conducted on a Li-Cor Odyssey infrared imaging system. GLT1 bands were quantified with ImageJ software by integrating band intensity followed by normalization to actin (Lepore et al., 2011c).

Functional glutamate uptake assay

Glia-induced precursor (GRP) cells were cultured in differentiation medium consisting of GRP basal medium supplemented with 10 ng/ml BMP4 and lacking the mitogen bFGF (Davies et al., 2008) for 2 d and then infected with AAV-Gfa2-eGFP or AAV-Gfa2-GLT1. After 6 d in culture, GRP medium consisting of GRP basal medium supplemented with 10 ng/ml BMP4 and lacking the mitogen bFGF (Davies et al., 2008) for 2 d and then infected with AAV-Gfa2-eGFP or AAV-Gfa2-GLT1. After 6 d in culture, GRP medium consisting of GRP basal medium supplemented with 10 ng/ml BMP4 and lacking the mitogen bFGF (Davies et al., 2008). Following infection, GRP-derived astrocytes were measured, as previously described (Dowd and Robinson, 1996), with slight modification. Briefly, cells were washed and preincubated with either a sodium-containing or choline-containing uptake buffer (containing (in mM): 35 Tris, 10 HEPES, 150 NaCl or choline chloride, 2.5 KCl, 1.2 CaCl₂, 1.2 MgCl₂, 1.2 K₂HPO₄, 10 glucose) for 20 min at 37°C and in dihydrokainate (DHK) treatment groups, 100 μM DHK was added to inhibit GLT1. The uptake buffer was then replaced with fresh uptake buffer containing 20 mM [14C]-glutamate (49 Ci/mmol; PerkinElmer) and 2 μM unlabeled glutamate. The cells were incubated for 5 min at 37°C. The reaction was terminated by washing cells three times with ice-cold uptake buffer containing 2 mM unlabeled glutamate,
followed by immediate lysis in ice-cold 0.1N NaOH. Cell extracts were then measured with a liquid scintillation counter (Beckman Instruments). The protein content in each well was measured using the Bradford protein assay (Bio-Rad).

In vitro scratch-wound assay
GLT1-overexpressing GRPs and wild-type GRPs, which both expressed GFP, were induced to differentiate into astrocytes with 10 ng/ml BMP-4. Astrocytes were cultured until confluent (10–14 d after plating). Scratch wounds were made as described previously (Nishio et al., 2005). Briefly, confluent cultures were scratched with a plastic pipette tip according to a preset grid to induce traumatic injury along the scratch wound. Culture dishes were washed twice with fresh DMEM to remove the debris and incubated with fresh medium for 24 h. The cells were then fixed, and immunostaining was performed with rabbit anti-Ki67 (1:200; Thermo Fisher Scientific) and mouse anti-GFAP (1:200; EMD Millipore) antibodies. Individual images (5–7 per scratch wound) were obtained at 10× magnification and then analyzed for cell-free area (in mm² per field), migrating cell numbers at various distances (D < 150 μm or D > 150 μm), proliferating cell numbers surrounding the wound site (using Ki67 as the proliferation marker), and lengths of migrating-cell processes (visualized with the GFP tag).

Cylinder paw preference test
All behavioral measurements were conducted on rats starting 2 weeks before injury and thereafter on a weekly basis until they were killed. Rats were weekly video-recorded while rearing freely in a transparent cylinder for 5 min. This test uses spontaneous rearing and exploratory behavior to test forelimb asymmetries under a preference paradigm. Three behaviors were scored during vertical exploration: use of the right ipsilateral forelimb-alone, use of the left contralateral forelimb-alone, and use of both forelimbs simultaneously. The hit score was expressed as a percentage of usage of each forelimb to total limb usage (Nicaise et al., 2012a).

Grip strength test
Muscle grip strength was weekly determined for the ipsilateral (right) forelimb using a "grip strength meter" (DFIS-2 Series Digital Force Gauge; Columbus Instruments). Grip strength testing was performed by allowing the rat to grasp a thin bar attached to the force gauge. This was followed by pulling the animal away from the gauge until the forelimb released the bar. The force measurements were averaged from three trials (Nicaise et al., 2012a).

Diaphragm compound muscular action potentials and spontaneous EMG recordings
Rats were anesthetized in the same manner described above. Phrenic nerve conduction studies were performed with single stimulation (0.5 ms duration; 6 mV amplitude) at the neck via near nerve needle electrodes placed along the phrenic nerve (Lepore et al., 2010). The ground needle electrode was placed in the tail, and the reference electrode was placed subcutaneously in the right abdominal region. Recording was obtained via a surface strip along the costal margin of the diaphragm, and compound muscular action potential (CMAP) amplitude was measured baseline to peak. Recordings were made using an ADI Powerlab 8/30 stimulator and BioAMP amplifier (ADInstruments), followed by computer-assisted data analysis (Scope 3.5.6, ADInstruments). For each animal, 10 tracings were averaged to ensure reproducibility.

Before being killed, animals were anesthetized and received a laparotomy. Bipolar electrodes 3 mm apart were inserted into several locations in the right hemidiaphragm (Nicaise et al., 2012a). The EMG signal was amplified, filtered through a bandpass filter (50–3000 Hz), and integrated (dV × dt) using LabChart 7 software (AD Instruments). Parameters, such as respiratory bursts per minute, burst duration, and integrated EMG peak amplitude, were averaged over 2 min sample periods.

Statistics
Results were expressed as means ± SEM. A Kolmogorov–Smirnov test was conducted for all variables to assess normality. Unpaired t test or Mann–Whitney was used to assess statistical significance between two groups. With respect to multiple comparisons involving ≥3 groups, statistical significance was assessed by one-way ANOVA followed by post hoc test (Bonferroni’s method). Statistics were computed with Graphpad Prism 5 (GraphPad Software). p < 0.05 was considered as statistically significant.

Results
Cervical contusion SCI resulted in both early and chronic downregulation of GLT1 promoter activity and protein expression
Using BAC-GLT1-eGFP transgenic promoter reporter mice, we spatially and temporally characterized the promoter activity of astrocyte GLT1 following unilateral cervical contusion SCI. In uninjured control mice, GLT1 promoter activity, as detected by eGFP + cells, was abundantly observed in gray matter and to a lesser extent in white matter (Fig. I A). At 2 DPI, confused spinal cords showed a disruption of spinal cord architecture, including significant loss of eGFP + astrocytes at the lesion epicenter (Fig. I B), compared with uninjured animals (Fig. I A). Higher-magnification views (Fig. I D, E) of the ventral horn were used to quantify individual eGFP + cells at specific distances from the epicenter, as illustrated in Figure 1C. We confirmed that a significant decrease in GLT1 promoter activity occurred as early as 2 DPI in sham versus injured mice (p < 0.001) and persisted to 14 and 42 DPI (p < 0.001) at the epicenter (Fig. I F). We next analyzed spinal cord segments rostral (Fig. I G, H) and caudal to the epicenter, locations where overt tissue disruption was not observed. In spinal cord ipsilateral to the lesion, the numbers of eGFP + cells 1.5 mm both rostral and caudal to epicenter were also significantly lower at 2 DPI (Fig. I H; p < 0.05). Although significant at only the later time point of 42 DPI (Fig. I L; p < 0.05), there were trends toward decreased numbers of eGFP + cells at 1.5 mm both rostrally at 14 DPI and caudally at 42 DPI compared with time-matched uninjured controls. When analyzed at the greater distance of 3.0 mm rostral and caudal to the epicenter, the numbers of eGFP + cells did not differ statistically between injured and uninjured mice at any time point (Fig. I J).

To complement these histological studies, we also correlated the downregulation of promoter activity with intraspinal GLT1 protein expression levels using immunoblotting on whole-tissue homogenate subdissected from various spinal cord segments (Fig. I C), GLT1 protein was highly expressed in uninjured spinal cord, while its expression was significantly reduced following cervical contusion (Fig. I K). At the epicenter, quantification of GLT1 immunoblotting showed ~50–80% downregulation at 2–42 DPI compared with uninjured control (Fig. I L). GLT1 protein expression at a spinal cord segment (C6) caudal to the epicenter was not different when comparing injured and uninjured animals (Fig. I M). Collectively, these results demonstrate that the numbers of GLT1-expressing astrocytes in ventral horn and overall intraspinal GLT1 protein expression were reduced soon after C4-contusion SCI injury and that the decrease persisted chronically for ≥6 weeks.

Lack of GLT1 expression in proliferating, reactive astrocytes following cervical contusion SCI
We next examined changes in the phenotype of astrocytes using markers of astrocyte reactivity (GFAP) and cell proliferation (Ki67) after cervical contusion SCI. Again using BAC-GLT1-eGFP transgenic mice, we correlated these immunohistochemical findings with GLT1 promoter activity. Astrocyte reactivity caused by injury results in robustly increased GFAP expression. In the intact spinal cord, gray matter astrocytes express levels of GFAP protein that are nearly undetectable using immunohistochemistry; therefore, GFAP analysis is an established approach
for histologically quantifying astrocyte reactivity at specific anatomical locations. As demonstrated by increased GFAP immunofluorescence intensity, astrocytes displayed dramatic reactivity in the ipsilateral ventral horn starting at 14 and 2 DPI at the lesion epicenter (Fig. 2A, B) and in intact rostral spinal cord (Fig. 2C, D), respectively, and reactivity persisted to 42 DPI. This reactive astrocyte response was associated with marked increases in the numbers of actively proliferating Ki67 cells in the ventral horn at both the lesion epicenter (Fig. 2A, B) and rostral spinal cord (Fig. 2C, D) at multiple time points postinjury.

We conducted confocal analysis to characterize changes in the phenotype of individual astrocytes following injury in the context of GLT1 expression and proliferation. In uninjured BAC-GLT1-eGFP cervical spinal cord, nearly all GFAP-positive astrocytes expressed GLT1 (Fig. 2E), which coincides with the robust and mostly astrocyte-specific expression of GLT1 in the CNS (Regan et al., 2007). Following cervical-contusion SCI, there was a pronounced reactivity of astrocytes in the ventral horn (both at the lesion epicenter and in intact rostral spinal cord) at all time points characterized by significantly increased expression of GFAP in individual astrocytes, as well as an increase in the total numbers of astrocytes (Fig. 2F). Following injury, there was also a large increase in the number of total proliferating cells in the ventral horn at epicenter and rostral locations, as assessed with Ki67 (Fig. 2F). A significant percentage of these Ki67+ cells were GFAP+ cells (Fig. 2F), suggesting that astrocyte proliferation at least partially accounted for the increase in number of astrocytes postinjury. Importantly, despite this increase in the number of GFAP+ cells, the majority of reactive astrocytes following injury (proliferating astrocytes and/or nondividing astrocytes that have increased GFAP expression) did not express GLT1 (Fig. 2F).

Collectively, these results show that astrocytes of the ventral horn become reactive very early following cervical contusion SCI and remain chronically in a reactive state. In addition, a large portion of the astrocyte population proliferates, particularly with proximity to the site of injury. Despite this robust response that can result in an increase in the number of astrocytes, these reactive cells were compromised in GLT1 expression following cervical SCI, suggesting an inability to properly regulate extracellular glutamate homeostasis.

Intraspinal AAV-mediated gene delivery resulted in overexpression of normally functioning GLT1 without inducing motor neuron loss or phenotypic impairment

To investigate whether overexpression of exogenous GLT1 in astrocytes protects PhMNs against glutamate excitotoxicity following cervical-contusion (C4) SCI in rats, we intraspinally delivered the GLT1-His fusion gene or eGFP gene using AAV-Gfa2
injured spinal cord, and found robust expression of His-tagged GLT1 protein in the ventral horn at day 10 postinjection, but little-to-no expression at day 2 (Fig. 3B). We injected AAV-Gfa2-eGFP or AAV-Gfa2-GLT1 into the injured spinal cord 10 d before unilateral C4 contusion, and we then examined the efficiency and location of transduction by the AAV-Gfa2 vector by detecting eGFP reporter distribution in the cervical spinal cord at day 10 postinjury. Expression of exogenous gene delivered by AAV-Gfa2 vector was highly localized to ventral gray matter, with significant transduction also observed in white matter close to the gray-white matter interface (Fig. 3B, C). The majority of eGFP-labeled cells possessed an astrocyte-like morphology (Fig. 3C, inset). eGFP+ cell counting showed that transduced cells were robustly distributed from −3 mm rostral to 3 mm caudal to a single injection site in both gray and white matter (Fig. 3D).

Although previous reports have shown that exogenous gene expression from AAV is nontoxic (Ponnazhagan et al., 1997; Monahan et al., 2002), we tested potential in vivo cytotoxicity in our experimental paradigm. AAV-Gfa2-GFP or AAV-Gfa2-GLT1 (10^13 genomic particles/ml), or artificial CSF as the control, was injected into ventral horn of the C4 spinal cord of uninjured rats. At day 10 postinjection, we conducted total motor neuron counts and phrenic nerve-diaphragm CMAP recordings, an in vivo electrophysiological assessment of functional diaphragm innervation by PhMNs (Lepore et al., 2010). There were no significant differences among the three groups in either ventral horn motor neuron numbers across the cervical spinal cord (Fig. 3E) or peak CMAP amplitudes (Fig. 3F), demonstrating that AAV injection itself and AAV-based GLT1 overexpression are not toxic in the spinal cord.

To assess the functional glutamate uptake capability of astrocytes transduced with virus, we performed a 3H-glutamate uptake assay on cultured astrocytes derived from glial-restricted precursors (a class of lineage-restricted astrocyte progenitor cells; Lepore et al., 2006), which were infected with AAV-Gfa2-eGFP or AAV-Gfa2-GLT1. Nearly all cells in both culture conditions were differentiated GFAP+ astrocytes (Fig. 3G). GLT1 protein expression was not detected in AAV-Gfa2-eGFP infected cells, which is expected in astrocytes cultured without neurons also present (Yang et al., 2009), while GLT1 was highly expressed in nearly all AAV-Gfa2-GLT1 astrocytes (Fig. 3G). Furthermore, GLT1 overexpression following AAV-Gfa2-GLT1 transduction significantly increased levels of sodium-dependent glutamate uptake in astrocytes (121.6±6.4 vs 190.2±24.4 cpm/μg protein, p<0.01), which could be blocked by 100 μM of the GLT1 inhibitor, DHK (p<0.05; Fig. 3H).

Collectively, these results demonstrate that AAV delivery resulted in efficient in vivo transduction and expression of GLT1 protein throughout the cervical spinal cord, did not result in neuronal injury or functional diaphragm impairment, and increased the capacity of astrocytes for functional GLT1-mediated glutamate uptake.

Intraspinal AAV-mediated gene delivery resulted in exogenous GLT1 overexpression specifically in astrocytes

We performed double immunostaining for GLT1 and His-tag, which is fused to the N terminus of GLT1, to distinguish exogenous from endogenous GLT1 protein in the injured spinal cord. His-tag colocalized with GLT1 (Fig. 4A, inset), and the His-tag+ area showed greater GLT1 immunofluorescence signal than nearby locations containing only endogenous GLT1 expression (Fig. 4A, arrowheads). Immunoblotting of whole-tissue spinal cord homogenate confirmed that total GLT1 protein expression levels in the injured spinal cord were significantly increased in the

 vectors that drive expression under control of the astrocyte GFAP promoter (Fig. 3A).

We first temporally assessed the time course of exogenous His-GLT1 expression at days 2 and 10 postinjection into the un-
AAV-Gfa2-GLT1 group compared with AAV-Gfa2-eGFP control ($p < 0.05$; Fig. 4B). Double immunostaining for GFAP and His-tag demonstrated that exogenous GLT1 was mainly expressed in GFAP+ astrocytes (arrowheads denote double-labeled cells). To further quantify the phenotypic identity of transduced cells in the spinal cord, we performed immunostaining for the lineage-specific markers GFAP (astrocytes), Olig2 (oligodendrocyte lineage), and NeuN (neurons) in both the uninjured (Fig. 4D) and injured (images not shown) spinal cord at day 10 following AAV-Gfa2-eGFP injection. Quantification results show that most eGFP+ cells were GFAP+ astrocytes (62.8 ± 3.4% of all GFP+ cells in uninjured tissue, 91.5 ± 1.0% in injured tissue), while only a small percentage of eGFP+ cells were cells of the oligodendrocyte lineage (18.5 ± 0.8% in uninjured tissue, 6.2 ± 0.5% in injured tissue) or neurons (4.2 ± 2.0% in uninjured tissue, 3.5 ± 0.7% in injured tissue; Fig. 4E). Interestingly, expression in astrocytes was especially pronounced in the injured spinal cord (i.e., 91.5% of transduced cells and greater eGFP intensity), likely due to the intense activation of the Gfa2 promoter following CNS trauma. High-magnification confocal images further show eGFP colocalization with GFAP, but not with Olig2 or NeuN (Fig. 4F). These results demonstrate that exogenous gene delivered by AAV-Gfa2 vector was specifically expressed in astrocytes in our SCI animal model and that AAV-Gfa2-GLT1 injection significantly increased the intraspinal expression of GLT1 protein in the cervical ventral horn.

Intraspinal GLT1 overexpression in astrocytes increased PhMN loss following cervical-contusion SCI

Given that astrocyte GLT1 overexpression could be successfully achieved via intraspinal AAV delivery, we sought to determine whether GLT1 overexpression would exert beneficial effects on lesion size and motor neuron loss. We injected AAV-Gfa2-eGFP or AAV-Gfa2-GLT1 into rats 10 d before C4 contusion (Figs. 3A,
Intraspinal GLT1 overexpression in astrocytes increased PhMN loss following cervical-contusion SCI. A, AAV-Gfa2-eGFP or AAV-Gfa2-GLT1 was intraspinally injected into rats 10 d before C4-contusion SCI. B, C, Representative images of cresyl violet staining illustrate the lesion epicenter in AAV-Gfa2-eGFP (B) and AAV-Gfa2-GLT1 (C) groups. D, E, Lesion size (D), rostral–caudal distance of lesion spread (E), and total lesion volume (F) were increased in AAV-Gfa2-GLT1 rats compared with AAV-Gfa2-eGFP animals. Neural populations in the ipsilateral ventral horn were identified (C, inset), quantified, and plotted at multiple distances from the epicenter (I, J). Representative images show longitudinal (I, K) and transverse (J, L) sections of the cervical spinal cord from AAV-Gfa2-eGFP and AAV-Gfa2-GLT1 pretreated rats, revealing ipsilateral labeling of PhMNs by injecting CTβ into the ipsilateral hemidiaphragm. This approach confirmed that AAV gene delivery spatially targeted astrocytes surrounding CTβ-labeled PhMNs (Fig. 5H). We analyzed CTβ labeling in both longitudinal (Fig. 5I, K) and transverse (Fig. 5J, L) spinal cord sections. Increased rostral–caudal extent of CTβ⁺ PhMNs. Injury epicenter is denoted by white asterisk.

Figure 5. Intraspinal GLT1 overexpression in astrocytes increased PhMN loss following cervical-contusion SCI. A, AAV-Gfa2-eGFP or AAV-Gfa2-GLT1 was intraspinally injected into rats 10 d before C4-contusion SCI. B, C, Representative images of cresyl violet staining illustrate the lesion epicenter in AAV-Gfa2-eGFP (B) and AAV-Gfa2-GLT1 (C) groups. D–E, Lesion size (D), rostral–caudal distance of lesion spread (E), and total lesion volume (F) were increased in AAV-Gfa2-GLT1 rats compared with AAV-Gfa2-eGFP animals. Neural populations in the ipsilateral ventral horn were identified (C, inset), quantified, and plotted at multiple distances from the epicenter (I, J). Representative images show longitudinal (I, K) and transverse (J, L) sections of the cervical spinal cord from AAV-Gfa2-eGFP and AAV-Gfa2-GLT1 pretreated rats, revealing ipsilateral labeling of PhMNs by injecting CTβ into the ipsilateral hemidiaphragm. This approach confirmed that AAV gene delivery spatially targeted astrocytes surrounding CTβ⁺ PhMNs (Fig. 5H). We analyzed CTβ labeling in both longitudinal (Fig. 5I, K) and transverse (Fig. 5J, L) spinal cord sections. Increased rostral–caudal extent of CTβ⁺ PhMNs. Injury epicenter is denoted by white asterisk. Loss of CTβ-labeled cells was greater at the rostral C3 (I–L) and caudal C5 (data not shown) boundaries of the lesion in the AAV-Gfa2-GLT1 group (K, L) compared with the AAV-Gfa2-eGFP group (I, J). M, Quantification of CTβ⁺-labeled cells shows significant loss of PhMNs at multiple distances from the epicenter. N, Increased loss of total CTβ⁺-labeled PhMNs was observed in the AAV-Gfa2-GLT1 group compared with the AAV-Gfa2-eGFP group. Results were expressed as means ± SEM. *p < 0.05; **p < 0.01; ***p < 0.001; AAV-Gfa2-GLT1 group versus AAV-Gfa2-eGFP group. n = 11 rats per group for lesion size analysis and total motor neuron counting; n = 4 rats per group for CTβ⁺ PhMN counting.
PhMN loss was observed in the AAV-Gfa2-GLT1 group (Fig. 5K), with the rostral population of PhMNs located at C3–C4 being particularly affected. As previously reported in rats (Mantilla et al., 2009; Nicaise et al., 2012a), PhMNs are concentrated within compact groups of cells forming a linear column throughout the ventromedial region of the C3–C5 ventral horn. Although both AAV groups showed similar PhMN loss at the epicenter, increased survival of CTB + cells was found in the most rostral (Fig. 5I,L, white arrowheads) and caudal (data not shown) aspects of the cervical region in AAV-Gfa2-eGFP rats. Only sparse CTB + cells were observed in the rostral region of AAV-Gfa2-GLT1 rats (Fig. 5L). Quantification at multiple distances from the lesion epicenter further demonstrated greater PhMN loss in the AAV-Gfa2-GLT1 group compared with AAV-Gfa2-eGFP (Fig. 5M). Increased PhMN degeneration was also quantified by estimating the total number of CTB + PhMNs across the C3–C5 region (ipsilateral to the injury) in the two AAV-injected groups (p < 0.05; Fig. 5N).

In summary, these histological results demonstrate that GLT1 overexpression in astrocytes of the ventral horn increased lesion size, total motor neuron loss, and specifically PhMN loss following cervical contusion SCI.

**Figure 6.** Intraspinal GLT1 overexpression worsened phrenic nerve axonal degeneration and diaphragm denervation. A, Phrenic nerve axonal degeneration was observed in both AAV-Gfa2-eGFP and AAV-Gfa2-GLT1 pretreated rats. B, Quantification of axonal density revealed a decreased number of intact fibers in the AAV-Gfa2-GLT1 group compared with the AAV-Gfa2-eGFP group. Diaphragm NMJs were assessed via labeling with Alexa-647-conjugated α-bungarotoxin (red), SMI-312R (green), and SV2-s (green). C–E, While areas of muscle denervation were observed in both AAV-Gfa2-eGFP (C) and AAV-Gfa2-GLT1 (D, E) groups, reduced numbers of intact NMJs and more widespread areas of denervated NMJs were detected in the AAV-Gfa2-GLT1 group. F–H, NMJ morphology analysis was divided into a number of established phenotypic categories: intact (F, asterisk), fully denervated (G, arrow), partially denervated (F, G, arrowheads), multiply innervated (H, double asterisk), and preterminal axon thinning. I, For analysis, the hemidiaphragm was divided into three anatomical regions (ventral, medial, and dorsal). J, Ipsilateral hemidiaphragm from the AAV-Gfa2-eGFP group showed milder NMJ abnormalities. K, In the AAV-Gfa2-GLT1-injected rats, we observed a reduced percentage of intact NMJs compared with AAV-Gfa2-eGFP-injected rats, regardless of the area examined. D, E, K, Compared with the ventral portion of the muscle in the AAV-Gfa2-GLT1 group (D), medial and dorsal locations in these same AAV-Gfa2-GLT1 animals were particularly affected (E), displaying a greater percentage of fully denervated NMJs (K). L, M, The percentages of NMJs exhibiting partial denervation (L) or multiple innervation (M) were similar between the two groups. N, The percentage of NMJs with signs of preterminal axon thinning was significantly reduced in the AAV-Gfa2-GLT1 group only at the dorsal region. Results were expressed as means ± SEM. *p < 0.05, **p < 0.01, ***p < 0.001 AAV-Gfa2-GLT1 group versus AAV-Gfa2-eGFP group. n = 11 rats per group for phrenic nerve analysis; n = 5 rats per group for NMJ analysis.

**Intraspinal GLT1 overexpression worsened phrenic nerve axonal degeneration and diaphragm denervation**

We next attempted to correlate central changes observed in the cervical spinal cord with peripheral pathology at the level of both the phrenic nerve and diaphragm NMJ. Anterograde degeneration in the phrenic nerve was more pronounced in the AAV-Gfa2-GLT1 group than in the AAV-Gfa2-eGFP group (Fig. 6A). Quantification of axon density confirmed a significant exacerbation of motor axon loss with AAV-Gfa2-GLT1 (Fig. 6B). To determine whether greater PhMN loss resulted in pathological alterations at the diaphragm NMJ, hemidiaphragm muscle was examined. While areas of muscle denervation were observed in both AAV-Gfa2-eGFP (Fig. 6C) and AAV-Gfa2-GLT1 (Fig. 6D,E) groups, reduced numbers of intact NMJs and more widespread areas of denervated NMJs were detected in the AAV-Gfa2-GLT1 group. To specifically characterize the types of pathological changes occurring, NMJ morphology analysis was divided into a number of established phenotypic cate-
Intraspinal GLT1 overexpression worsened both diaphragmatic and forelimb motor dysfunction

It is important to characterize the in vivo functional effects of AAV-based GLT1 overexpression on both forelimb motor and diaphragmatic function, given that the motor neuron populations innervating these muscles are located in the cervical spinal cord and are profoundly affected by human cervical SCI. We monitored forelimb motor function after cervical contusion weekly for 6 weeks postinjury using grip strength testing and the paw preference/cylinder test. While both AAV groups were chronically impaired up to 42 DPI compared with preinjury baseline, AAV-Gfa2-eGFP animals showed signs of partial grip strength recovery at several time points postinjury compared with AAV-Gfa2-GLT1 rats (p < 0.05 at 21, 28, and 42 DPI; Fig. 7A). This finding was accompanied by an increased use of ipsilateral forelimb in the cylinder test in the AAV-Gfa2-eGFP group (p < 0.05 at 21 and 35 DPI, p < 0.01 at 42 DPI; Fig. 7B).

In parallel, we electrophysiologically investigated functional innervation of the diaphragm by PhMNsin using phrenic nerve conduction studies. Following supramaximal phrenic nerve stimulus, we obtained CMAP recordings from the surface of the ipsilateral hemidiaphragm. In both groups, peak CMAP amplitudes were significantly reduced with AAV-Gfa2-GLT1 injection at 2 (D) and 14 (E) DPI. F, G, Spontaneous diaphragm EMG recordings from AAV-Gfa2-eGFP (F) and AAV-Gfa2-GLT1 (G) groups are shown as raw voltage output (F, G, top) and integrated signals (F, G, bottom). H-J, Inspiratory burst duration (H), inspiratory bursts per minute (I), and integrated EMG amplitude (J) in the ipsilateral hemidiaphragm were averaged over a 2 min sample period. Results were expressed as means ± SEM. *p < 0.05, **p < 0.01 AAV-Gfa2-GLT1 group versus AAV-Gfa2-eGFP group. n = 11 rats per group for assessment of forelimb motor function; n = 5–7 rats per group for CMAP recordings; n = 4 rats per group for EMG recordings.
Intraspinal Gfα2-GLT1 gene delivery at the time of cervical-contusion SCI did not affect histological or functional outcomes. AAV-Gfα2-eGFP or AAV-Gfα2-GLT1 vector was intraspinally injected into rats immediately following C4 contusion. Animals were killed 35 DPI. B–F, Lesion size (B), rostral–caudal extension of the lesion (C), total lesion volume (D), ipsilateral total motor neuron counts (E), and axon density in the ipsilateral phrenic nerve (F) did not significantly differ between groups. G, H, No differences were noted between groups with grip strength testing (G) or the cylinder paw preference test (H). I, Diaphragm CMAP amplitudes did not differ between groups. Results were expressed as mean ± SEM. *p < 0.05, **p < 0.01 AAV-Gfα2-GLT1 group versus AAV-Gfα2-eGFP group. n = 5 rats per group for lesion size analysis and total motor neuron counting; n = 9 rats per group for phrenic nerve analysis, assessment of forelimb motor function, and CMAP recordings.

Overexpression of GLT1 reduced astrocyte proliferation and migration, altered astrocyte morphology in vitro, and compromised astroglial scar border formation in vivo

To determine the phenotypic effects of GLT1 overexpression on astrocytes, we performed an in vitro scratch-wound assay and also analyzed astroglial scar formation in vivo following cervical contusion SCI to study the protective scar-forming capacity of astrocytes in response to injury. For the in vitro scratch-wound assay, wound/gap area, astrocyte migration to various distances within the scratch site (D < 150 μm or D > 150 μm), cell proliferation surrounding the wound site, and process length of migrating cells were assessed 24 h after scratch injury to an astrocyte monolayer was induced (Fig. 9A). Quantification results showed that overexpression of GLT1 using AAV-Gfα2-GLT1 slowed wound healing (Fig. 9B), reduced astrocyte migration (Fig. 9C) and proliferation (Fig. 9D), and shortened the process of migrating astrocytes (Fig. 9E). For the in vivo astroglial scar border formation analysis, immunostaining with anti-GFAP antibody

of each discharge (Fig. 7H), while the number of inspiratory bursts per minute was significantly increased in the AAV-Gfα2-GLT1 group compared with control group (Fig. 7J). The integrated EMG amplitude in the AAV-Gfα2-GLT1 group was slightly reduced compared with the control group; however, this difference was not statistically significant (Fig. 7J).

These results demonstrate that increased cervical motor neuron loss and diaphragm denervation associated with astrocyte GLT1 overexpression resulted in an actual in vivo worsening of diaphragm and forelimb motor dysfunction in our clinically relevant model of cervical-contusion SCI.

Intraspinal GLT1 gene delivery at the time of cervical-contusion SCI did not affect histological or functional outcome

We next sought to determine the effects of AAV-Gfα2-GLT1 gene delivery at an “acute” time point by injecting virus immediately following unilateral cervical contusion using the exact same SCI model (Fig. 8A). AAV pretreatment represents a powerful approach for testing proof of principle because it allows for maximal GLT1 overexpression by the time of injury, even if it is not directly relevant to clinical translation. In contrast, the acute injection strategy is a delivery paradigm directly relevant to the potential treatment of cervical SCI patients. Using the same histological analyses employed for the AAV pretreatment studies, we characterized the effects of AAV-Gfα2-eGFP and AAV-Gfα2-GLT1 following acute injection. There were no significant differences between virus types in lesion size (Fig. 8B), rostral-caudal extension of the lesion (Fig. 8C), lesion volume (Fig. 8D), total motor neuron counts (Fig. 8E), or phrenic nerve axonal loss (Fig. 8F) at 35 DPI. Given the lack of differences in a number of histological tests, we did not conduct CTB or NMJ analyses on these animals. We also conducted functional examination of forelimb motor and diaphragmatic function in the acute injection paradigm. Similar to histological analyses, no differences between groups were noted in ipsilateral forelimb grip strength (Fig. 8G), ipsilateral forelimb use in the cylinder test (Fig. 8H), or diaphragm CMAP amplitude at 35 DPI (Fig. 8I).

Last, we also tested delayed injection of AAV vectors at 7 DPI, again using the exact same SCI model as pretreatment and “acute” injection paradigms. With delayed injection of AAV-Gfα2-eGFP and AAV-Gfα2-GLT1, we observed no effects of GLT1 overexpression on lesion measurements, total motor neuron or PhMN loss, diaphragm denervation, or forelimb motor function using histological and functional measures (data not shown).

Unlike the results obtained with pretreatment, our findings with “acute” injection and injection at 7 DPI demonstrate that AAV-GLT1 delivery soon after injury does not modify outcome. These results are likely due to the temporal delay involved in achieving transduction and GLT1 overexpression with intraspinal AAV injection (Fig. 3B). If GLT1 overexpression is to be targeted at early time points post-SCI, an alternative strategy to viral vector delivery is likely needed; however, intraspinal injection immediately after injury (or even at very early time points) will likely not be a practically feasible approach for patients. Last, the difference in results between pretreatment and acute/delayed delivery paradigms suggests that the negative effects of GLT1 overexpression are based on events occurring during the initial hours to days post-trauma.
was performed 3 weeks after injury (Fig. 9F) in the pretreatment virus delivery paradigm. Quantification results show that astroglial scar border formation (Fig. 9F, yellow solid line) in the AAV-Gfa2-GLT1 group was significantly reduced compared with the AAV-Gfa2-eGFP group (Fig. 9G). In addition, GFAP signal intensity of the astrocyte scar border (Fig. 9F, red selection, a') in the AAV-Gfa2-GLT1 group was significantly reduced compared with the AAV-Gfa2-eGFP group, while in the area next to the border (Fig. 9F, red selection, a) GFAP signal intensity was similar between the two groups. Results were expressed as means ± SEM. *p < 0.05, **p < 0.01, ***p < 0.001. n = 5–7 per group for in vitro assay, n = 12–13 per group for in vivo analysis.

Discussion
In the context of SCI, astrocyte reactivity has traditionally been viewed negatively in association with the astroglial scar and its inhibitory effects on axon regeneration/plasticity. However, astrocytes normally play an integral role in the CNS via a number of important functions (Pekny and Nilsson, 2005). The astrocyte response following SCI should not be approached only as a dichotomy of quiescent versus reactive, but should be viewed as occurring along a spectrum of phenotypes, depending on type, location, timing, and severity of the insult (Sofroniew, 2005). Importantly, it is crucial to study specific astrocyte functions in the diseased nervous system, not just to generalize an astrocyte response. Along these lines, we observed both early and persistent GLT1 expression loss in ventral horn astrocytes following cervical contusion. These findings are in line with previous work from...
the astrocyte population did not express GLT1. In our experiments, more GLT1 would be available for such a mechanism. This would be partly consistent with results of ischemic injury. At early stages (i.e., 5 min), ischemia-induced glutamate levels in hippocampus of wild-type mice are significantly lower than those in mice lacking GLT1, while after 20 min of ischemia, glutamate levels in wild-type mice are double those of GLT1 heterozygotes (Mitani and Tanaka, 2003). These results suggest that GLT1 forward transport plays a role in uptake at early stages, while reverse transport may subsequently predominate for some period of time, depending on specifics of the insult (Szatkowski and Attwell, 1994). Since neuron death caused by oxygen-glucose deprivation preconditioning is relieved by down-regulating (Wang et al., 2013) or inhibiting (Kosugi and Kawahara, 2006) GLT1, the combination of GLT1 overexpression and selective inhibition of reverse transport may be a successful approach. Unlike the commonly used GLT1 inhibitors DL-threo-β-benzylxystapartate (Colleoni et al., 2008) and DHK (Shirasaki et al., 2010), which target both forward and reverse transport, bromocriptine (Shirasaki et al., 2010) and (−3-hydroxy-4,5,6,6a-tetrahydro-3aH-pyrrolo[3,4-d]-isoxazole-4-carboxylic acid (Colleoni et al., 2008) are thought to mainly inhibit reverse transport.

Astrocytes are more resistant to ischemic injury than neurons (Goldberg and Choi, 1993), possibly because of larger glycerol stores (Cataldo and Broadwell, 1986; Brown and Ransom, 2007) and reduced susceptibility to excitotoxicity (Choi and Rothman, 1990). However, in severe ischemic conditions, loss of the Na+/K+ gradient across the astrocyte plasma membrane can result in astrocyte death in the presence of glutamate (Rossi et al., 2007; Szydlowska et al., 2010). It has been reported that GLT1 reverse transport promotes astrocyte survival by maintaining low intracellular concentrations of Na+ and Ca2+ (Kosugi and Kawahara, 2006). In our experiments, GLT1 overexpression may further promote survival of reactive astrocytes. However, chronic astrocyte reactivity may result in a phenotypic switch to a detrimental population. These highly reactive astrocytes could release nitric oxide and reactive oxygen species (Bal-Price and Brown, 2001) that modulate glutamate receptors and consequently lead to damage (Stewart et al., 2002). These cells may also release a variety of cytokines, including TNFα, that are toxic to oligodendrocytes and inhibit neurite outgrowth (D’Souza et al., 1996; Neumann et al., 2002). CCL2, CCL5, and CXCL18 release could also recruit immune cells, leading to inflammatory damage (Oh et al., 1999).

After CNS trauma, areas of focal damage are surrounded by an astroglial scar that separates necrotic from spared tissue (Kawano et al., 2012). Transgenic ablation of the proliferating astrocytes that contribute to this process post-SCI leads to increased inflammation, demyelination, lesion volume, and neuronal loss (Faulkner et al., 2004). The glial scar is formed, in part, by proliferating elongated astrocytes. Similarly, when cocultured in vitro with inflammatory cells, astrocytes display bipolar or radial morphologies with elongated processes, and segregate macrophages into discrete clusters (Wanner et al., 2013). In an in vitro...
scratch–wound experiment using astrocyte cultures, we found that GLT1 overexpression reduced astrocyte migration and proliferation, and altered astrocyte morphology such that they displayed shortened processes compared with cells transduced with control virus. Furthermore, in vivo scar formation analysis showed that astroglial scar border formation was significantly compromised in the AAV–GLT1 compared with the control AAV–eGFP group following cervical-contusion SCI. These findings suggest that alteration of the protective scar-forming capacity of astrocytes may be one possible mechanism for the detrimental effects of GLT1 overexpression.

Beneficial effects of GLT1 have been reported in other CNS disease models. For example, GLT1 overexpression reduces ischemic brain injury in a rat stroke model (Harvey et al., 2011). GLT1 knockdown exacerbates hippocampal neuronal damage following traumatic brain injury (Rao et al., 2001). Intraspinal gene transfer of GLT1 attenuates inflammatory and neuropathic pain in rats (Maeda et al., 2008). The discrepancy with our data may be explained, for example, by disease-specific pathological conditions and different spatiotemporal GLT1 expression patterns.

We did not detect motor neuron toxicity or functional impairment with AAV injection into uninjured spinal cord or in the “acute” or “delayed” paradigms postinjury. We found worsening in the pretreatment paradigm only with AAV–GLT1. We observed robust and mostly astrocyte-specific transduction that resulted in significantly increased intraspinal GLT1 expression, with very little ectopic expression in nonastrocyte populations. Last, we found increased functional glutamate uptake with AAV–GLT1, demonstrating that transduction resulted in not only protein overexpression but correct membrane localization and transporter function. Together, these results suggest that the observed negative effects were due to increased GLT1 expression in astrocytes and not to the injection procedure, AAV vector, or expression of nonfunctional transporter.

Despite the current results, we still believe that increasing GLT1 is a potentially useful approach. Further studies need to determine the target astrocyte population(s) and correct timing of delivery. Importantly, our findings demonstrate that this AAV-based strategy allows for efficient and long-term gene delivery to spinal cord astrocytes. Furthermore, this anatomically specific approach is crucial for avoiding unwanted side effects. Although AMPA and NMDA antagonists have shown promise for SCI (Wrathall et al., 1994; Agrawal and Fehlings, 1997; Rosenberg et al., 1999; Gaviria et al., 2000), therapeutically targeting glutamate receptor overactivation has diffuse and nonspecific effects in CNS, PNS, and potentially other tissues, as well as only a short temporal duration (Bleakman et al., 2006).

Studies need to model clinically relevant aspects of human SCI. Despite the predominance of cervical contusion in patients, therapies targeting protection of respiratory circuitry have not been extensively tested in appropriate models. The data presented illustrate how our cervical-contusion model and set of histological and functional analyses represents a powerful approach for understanding and treating SCI.

References


