Cbp-dependent histone acetylation mediates axon regeneration induced by environmental enrichment in rodent spinal cord injury models

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After a spinal cord injury, axons fail to regenerate in the adult mammalian central nervous system, leading to permanent deficits in sensory and motor functions. Increasing neuronal activity after an injury using electrical stimulation or rehabilitation can enhance neuronal plasticity and result in some degree of recovery; however, the underlying mechanisms remain poorly understood. We found that placing mice in an enriched environment before an injury enhanced the activity of proprioceptive dorsal root ganglion neurons, leading to a lasting increase in their regenerative potential. This effect was dependent on Creb-binding protein (Cbp)–mediated histone acetylation, which increased the expression of genes associated with the regenerative program. Intraperitoneal delivery of a small-molecule activator of Cbp at clinically relevant times promoted regeneration and sprouting of sensory and motor axons, as well as recovery of sensory and motor functions in both the mouse and rat model of spinal cord injury. Our findings showed that the increased regenerative capacity induced by enhancing neuronal activity is mediated by epigenetic reprogramming in rodent models of spinal cord injury. Understanding the mechanisms underlying activity-dependent neuronal plasticity led to the identification of potential molecular targets for improving recovery after spinal cord injury.

INTRODUCTION  
After spinal cord injury (SCI), motor and sensory axons fail to regenerate, leading to permanent neurological impairments (1, 2). This absence of regeneration in the central nervous system (CNS) after injury has been attributed to two main interconnected factors: the presence of growth inhibitory molecules in the CNS and the lack of an effective neuronal-intrinsic regenerative response (2–4). Although axon regeneration fails in the CNS after injury, limited regeneration and partial functional recovery do occur in the injured peripheral nervous system (PNS) (5, 6). Dorsal root ganglion (DRG) neurons are located at the interface between the PNS and CNS. Their pseudounipolar anatomy provides the unique opportunity to study regenerative mechanisms in both locations from a single cell body. A well-established model for increasing the intrinsic regenerative capacity of DRG neurons in rodents consists of inducing a conditioned injury to the axons in the peripheral sciatic nerve before axons in the CNS. The peripheral injury induces the expression of regeneration-associated genes (RAGs) that increases the regenerative potential of the DRG neurons eliciting a stronger regenerative response after CNS injury (7–9). Although this has provided useful insights in the mechanisms underlying axon regeneration, a conditioning lesion only induces modest regeneration in the CNS (9) and is not a clinically viable approach.

Sensory DRG neurons convey afferent information from the periphery to the spinal cord, to modulate motor outputs, and to supraspinal structures for the elaboration of sensorimotor integration and conscious perception. Among the DRGs, proprioceptive neurons innervate muscle spindles and Golgi tendon organs (10). They transmit information about the length and tension of muscles, which plays a critical role during sensorimotor execution (11, 12). Proprioceptive afferent feedback influences the activity of circuits in the spinal cord and plays an essential role in providing immediate adjustment and refinement of movement and motor learning (13). Moreover, it plays an important role in directing motor recovery after SCI. Mice lacking functional proprioceptive afferents exhibit a defective rearrangement of descending pathways that prevents recovery after SCI (12). This observation suggests that proprioceptive neurons may deliver molecular cues for axonal regrowth and sprouting after injury. In support of this hypothesis, previous studies have shown that modulation of proprioceptive afferents with electrical stimulation (14, 15) and rehabilitative training (16) augmented neuroplasticity and recovery in both animal models and humans after SCI (17). However, the underlying mechanisms remain poorly understood.
Here, we used environmental enrichment (EE) as a means to physiologically increase neuronal activity in rodents. It has been previously shown that exposing mice to EE has profound effects on the structure and function of cortical and hippocampal neurons, leading to enhanced neuroplasticity, synaptogenesis, and neurogenesis and ultimately improved memory and cognitive performance (18–20). The innervation of muscle spindles and Golgi tendon organs by proprioceptors suggests that they are ideally located to be modulated by environmental stimuli.

We investigated the impact of EE on proprioceptive DRG neurons, hypothesizing that this would prime them to initiate a regenerative response to a subsequent injury, similar to a conditioning injury. Our results showed that augmenting the activity of proprioceptive DRG neurons using EE in mice induced a lasting increase in their regenerative potential due to Creb (cyclic adenosine monophosphate–response element–binding protein)–binding protein (Cbp)–mediated histone acetylation. Pharmacologically increasing Cbp acetylation activity after SCI in a mouse model induced regeneration and sprouting of sensory axons and brainstem motor pathways, resulting in improvements in both sensory and motor functions. These results show that neuronal activity leads to changes in chromatin environment that boost the regenerative capacity of neurons. Elucidating the mechanisms underlying activity–dependent responses in neurons translated into the identification of a potential treatment for SCI that warrants evaluation in humans.

**RESULTS**

**EE induces a lasting increase in the regenerative potential of DRG neurons**

To test whether EE can enhance the regenerative potential of DRG neurons, mice were exposed to EE or standard housing (SH) for 1, 3, 6, 10, or 35 days. After exposure to EE, the sciatic DRGs were cultured in a growth-permissive substrate for 12 hours. Neurite outgrowth of DRG neurons was enhanced after exposure to EE for 10 or 35 days (Fig. 1A and B). The extent of neurite outgrowth on a growth inhibitory myelin substrate was similar to what is observed after a conditioning injury (Fig. S1).

The EE-dependent increase in neurite outgrowth was abolished when incubating the neurons with the transcriptional inhibitor actinomycin-D (fig. S2), suggesting a dependence of this response on gene transcription. EE-dependent increase in neurite outgrowth was maintained when the animals returned to SH for 5 weeks after 10 days of EE (Fig. 1, C to E), suggesting that EE may trigger long-lasting effects. Analyses of growth responses revealed that EE enhanced axon elongation rather than branching, which may translate into axonal regeneration in vivo (Fig. S3).

Given the multifactorial nature of EE, we decided to discriminate between the relative role of running, which has previously been shown to enhance peripheral nerve regeneration (21), and the remaining environmental stimuli (larger cage, increased number of mice, previously unidentified objects, and increased nesting material) on DRG outgrowth. Mice were placed either in EE, in SH, in EE with an immobilized wheel, or in SH with a running wheel for 10 days. Analysis of neurite outgrowth showed that, although the running wheel in SH enhanced outgrowth compared to SH alone, the full complement of EE still induced a higher degree of outgrowth (fig. S4). This result indicated that the full EE is required for maximal enhancement of DRG outgrowth.

We next tested whether EE can also enhance regeneration of axons within the PNS and the CNS. Exposing the mice to EE before sciatic nerve transection and reanastomosis enhanced regenerative growth (Fig. 1, F and G). The same protocol also increased muscle reinnervation after a sciatic nerve crush (fig. S5).

We then assessed whether preexposure to EE would enhance regeneration of sensory axons in the dorsal columns after an SCI and compared this response to the regeneration observed after a conditioning injury. Three groups of adult mice were exposed to EE or SH for 10 days and then housed in SH. Subsequently, SH mice received a conditioning sciatic nerve axotomy (SNA) or sham injury. The EE groups only received the sham injury. Next, all the mice received a thoracic (T12) dorsal hemisection. Five weeks later, the retrograde tracer cholera toxin subunit B (CTB) was injected into the sciatic nerves to evaluate the regeneration of ascending dorsal column axons (Fig. 1H). Most of the labeled axons from SH sham mice retracted from the injury site. In contrast, labeled axons from EE mice could be observed within the lesion epicenter. Some regenerating axons even expanded beyond the lesion site, attaining a distance of up to 800 μm from the lesion border (Fig. 1H). As expected, the conditioning injury in SH mice also promoted axonal regeneration (Fig. 1, H and I).

Six weeks after injury, we conducted terminal electrophysiological experiments to evaluate the functionality of these regenerating axons. The dorsal columns were stimulated below the injury at lumbar vertebra 5 (L5) and recorded both below and above the lesion at L1 and T9, respectively (Fig. 1J). The amplitude of the compound action potentials recorded above the lesion site was larger for the animals that had been housed in EE compared to SH or SH-SNA (Fig. 1, K and L, black traces), suggesting an increase in neuroplasticity across the lesion. There was no difference in the compound action potentials recorded below the lesion for any of the groups (Fig. 1K, blue traces).

To determine whether regenerating sensory axons were responsible for the increase in conduction through the lesion site, we selectively silenced these axons using designer receptors exclusively activated by designer drugs (DREADD), as described previously (22). We injected the adeno-associated viral vector (AAV) AAV-flex-hM4Di into the sciatic nerve, which will silence the activity of DRG neurons after Cre recombination. Three weeks after SCI, an AAV-Cre was injected rostral to the site of injury to induce the expression of the Cre-dependent Gc coupled DREADD receptor only in those DRG neurons that had extended axons through and beyond the lesion. After three additional weeks, mice previously exposed to EE underwent electrophysiological assessment. Chemogenetic-mediated silencing restricted to regenerated axons (Cre-dependent AAV-flex-hM4Di) reduced conduction across the lesion, establishing causality that was confirmed with a reanastomosis (Fig. 1, K and L).

**Proprioceptive afferent feedback is required for EE-mediated increase in DRG regenerative growth**

We next investigated whether a specific type of DRG neuron was implicated in EE-dependent regenerative growth. We injected CTB into the distal sciatic nerve 1 day after performing a sciatic nerve crush injury. Consequently, only DRG neurons that regenerated an axon across the injury site and into the denervated nerve would be able to take up the CTB tracer (Fig. 2A). Two days after the CTB
injection, we assessed the number of CTB-positive DRG neurons that stained for markers of the main DRG subpopulations (Fig. 2B). Previous exposure to EE increased the number of CTB-positive DRG neurons, confirming that EE enhances axon regeneration (Fig. 2C). Most of the DRG neurons, which regenerated axons through the sciatic nerve crush and were retrogradely labeled with CTB, expressed markers of proprioceptors [parvalbumin (PV)] rather than nociceptors (isolectin B4 or substance P; Fig. 2, B and D), suggesting that EE preferentially enhanced the regeneration of proprioceptive DRG neurons in this model.

To evaluate whether the EE-dependent increase in regenerative potential of proprioceptive neurons relied on a muscle spindle proprioceptive mechanism, we used Egr3−/− mice. This mutation abolishes muscle spindle proprioceptive feedback while retaining a similar number of PV DRG neurons compared to wild-type (WT) mice (Fig. 2E) (12, 23). We found that EE-dependent but
not conditioning injury–dependent DRG outgrowth was abolished in mice lacking intact muscle spindles (Egr3−/−) (Fig. 2, F and G). This observation indicates that EE-dependent regenerative priming is likely to be contingent on intact proprioceptive afferent feedback.

Last, to further confirm the cell type specificity of the EE mechanism, we assessed neurite outgrowth of DRG neurons from mice that expressed the fluorescent marker tdTomato under the control of the PV promoter. Exposure to EE increased the outgrowth of tdTomato-positive (PVON) but not of tdTomato-negative (PVOFF)
DRG neurons (Fig. 2, H and I). Together, these experiments demonstrate that previous exposure to EE primes proprioceptive DRG neurons for enhanced axon regeneration.

**EE-induced signaling pathways involved in neuronal activity, calcium mobilization, and the regenerative program of large-diameter DRG neurons**

EE and exercise have been shown to increase neurotrophin expression and modify cytokines influencing neuroplasticity (21, 24–26). Unexpectedly, we found no changes in neurotrophin or cytokine levels in the DRG or in the blood (fig. S7), suggesting that alternative mechanisms may be responsible for EE-dependent DRG regenerative growth.

To uncover EE-dependent molecular mechanisms, we performed RNA sequencing (RNAseq) from whole DRGs or laser-captured large-diameter DRG neurons (LDNs), which represent the proprioceptor and mechanoreceptor population, as well as conducting proteomic analysis from sciatic nerve axoplasm after 10 days in EE or SH. Unsupervised gene expression clustering showed marked changes in gene expression of LDNs after EE compared to SH but not in the whole DRGs, which contains a multitude of different neuronal and glial cell types (Fig. 3A, fig. S8A, and data file S1). The data show that EE preferentially increased rather than repressed gene expression in LDNs (fig. S8B). Together, these results confirmed that EE specifically modulates gene expression in proprioceptors and mechanoreceptors. Similarly, proteomic analysis showed a larger number of up-regulated proteins after EE versus SH (71 versus 49; fig. S8C and data file S1). Overall, 37 of 71 up-regulated proteins were also up-regulated at the RNA level, suggesting that gene transcription drove more than half of the observed protein changes (data file S1).

Functional classification of EE-dependent gene expression changes in LDNs revealed that EE strongly modulated functionally interconnected molecular pathways involving ion channels, neuronal activity, calcium signaling, energy metabolism, and neuronal projection (Fig. 3B, fig. S8D, and data file S2). Combined analysis of the RNAseq and proteomic datasets for protein–protein interactions identified multiple interactions between proteins involved in neuronal activity, calcium signaling, and cytoskeletal rearrangements, supporting a role for EE-mediated activity in axon projection and elongation (fig. S8E).

![RNA sequencing in DRG](image1)

**Fig. 3. EE induces signaling pathways involved in neuronal activity, calcium mobilization, and the regenerative program of LDNs.** (A) Heatmap of the differentially expressed (DE) genes in whole-DRGs and LDNs RNAseq (P < 0.05). Color scale represents arbitrary expression units (lowest, blue; highest, red). (B) Pie chart of genes in each functional group identified by Gene Ontology analysis of DE genes in LDNs. Functional groups are color coded. (C) Representative images of sciatic nerves transduced with AAV5-GFP (green fluorescent protein), AAV5-hM4Di-mCitrine, or AAV5-hM3Dq-mCitrine labeled with mCitrine/GFP after sciatic nerve crush. Arrowhead, lesion site. Scale bar, 500 μm. (D) Quantification of axon regeneration (mean ± SEM, two-way repeated measures ANOVA, Tukey’s post hoc, **P < 0.001; n = 3 per group). (E) Representative time-lapse images of intracellular calcium release from whole-mount PV-glycogenically encoded calmodulin protein (GCaMP) DRGs before and after addition of 150 mM KCl. Scale bar, 50 μm. (F) Quantification of F/Fl (fluorescence intensity relative to baseline) ratio after 50 mM, 100 mM, and 150 mM KCl (mean ± SEM, two-way ANOVA, Sidak’s post hoc, ***P < 0.01, **P < 0.001, respectively; n = 4 per group).

These results encouraged us to investigate the role of neuronal activity and calcium release in the EE-mediated increase in DRG regenerative potential. We used an AAV-mediated chemogenetics approach to inhibit or enhance neuronal activity of DRG neurons using DREADD technology. When activated by the pharmacologically inert ligand clozapine-N-oxide (CNO), Gq-coupled (hM4Di) DREADD receptors inhibit adenylyl cyclase, which silences neuronal activity (27). Alternatively, activation of Gq-coupled (hM3Dq) DREADD receptors enhances neuronal activity by eliciting inositol 1,4,5-trisphosphate–mediated calcium release from intracellular...
stores (27). This increase in intracellular calcium activates calcium-dependent signaling cascades, which would, if our hypothesis is correct, mimic the effects of EE on DRG neurons identified with gene and proteomic analysis.

We injected AAV vectors into the sciatic nerve to express hM4Di, hM3Dq, or GFP in DRG neurons. mCitrine/GFP expression confirmed an efficient transduction (fig. S9). Four weeks after injection, mice expressing hM4Di, hM3Dq, or GFP were placed in EE or SH. CNO was added in the drinking water to activate the receptors. Ten days later, we performed a sciatic nerve crush and assessed the extent of regeneration 3 days after injury (Fig. 3, C and D).

Gq activation enhanced axon regeneration in SH mice, and Gi-dependent inactivation of DRG neurons attenuated axon regeneration of mice exposed to EE, demonstrating the importance of neuronal activity for EE-mediated regeneration (Fig. 3, C and D). Expression of Gi in SH mice or Gq in EE mice did not further impair or promote axon regeneration (Fig. 3, C and D). Similar results were obtained when assessing neurite outgrowth of DRGs cultured for 12 hours after in vivo DREADD transduction and EE exposure (fig. S10). Gq expression in DRGs from SH mice increased neurite outgrowth to an extent similar to what was observed in the DRGs of mice exposed to EE, whereas the expression of Gi in DRG neurons reduced EE-dependent outgrowth (fig. S10).

RNAseq and proteomics data suggested that EE may enhance calcium signaling in proproceptive DRG neurons. To visualize the induction of activity-dependent signaling pathways in proproceptive neurons in response to EE, we directly measured intracellular calcium as an indicator of release from intracellular stores. Transgenic mice expressing the calcium indicator GCaMP under the PV promoter were exposed to EE or SH for 10 days. Whole sciatic DRGs were extracted to measure the fluorescent signals of proprioceptive neurons ex vivo when applying increasing levels of potassium chloride. As anticipated, we found that exposing mice to EE increased the amount of calcium in DRG neurons at all the potassium chloride concentrations tested compared to DRGs from SH mice (Fig. 3, E and F, and movies S1 and S2). These observations show that EE increases calcium signaling in proprioceptive neurons, likely through increased solicitation of muscle spindles and force sensors within the enriched environment. These results provide evidence that EE increases neuronal activity and calcium signaling in proprioceptive DRG neurons and plays an important role in the activity-dependent increase in regenerative potential.

**Pharmacological activation of Cbp/p300 promotes sensory axon regeneration and recovery after a dorsal hemisection SCI in mice**

The central role of Cbp suggested that the pharmacological activation of Cbp might mimic the EE-dependent increase in regenerative growth of DRG neurons. To test this hypothesis, we delivered a small-molecule activator (TTK21) of the closely related transcriptional coactivators Cbp and p300 after SCI (36). Recent studies have shown that TTK21 is nontoxic in animal models and, when conjugated to glucose-derived carbon nanospheres (CSP), successfully crosses the blood-brain barrier, effectively enhances histone acetylation in the hippocampus, and promotes improvements in learning and memory capacities (36, 37).

We first confirmed that the addition of CSP–TTK21 to DRG cultures was capable of increasing neurite outgrowth compared to control CSP. CSP–TTK21 increased neurite outgrowth and H4K8 acetylation and reduced neurite branching in cultured DRG neurons (Fig. 5, A and B). We next tested whether CSP–TTK21 could promote axonal regeneration in vivo after a midthoracic dorsal hemisection (Fig. 5C). Injured mice received a weekly intraperitoneal injection of CSP–TTK21 (20 mg/kg) or control CSP, beginning 4 hours after injury. To assess regeneration of sensory axons in the dorsal columns, we injected the sciatic nerves with CTB tracer and examined the spinal cord 6 weeks after SCI (Fig. 5D). Treatment with CSP–TTK21 promoted sensory axon regeneration of up to 1000 μm rostral to the lesion (Fig. 5, E and F). We conducted behavioral assessments for 5 weeks after SCI to evaluate recovery of sensorimotor function. We selected tasks contingent on accurate proprioceptive information, which included the gridwalk and adhesive tape test. We observed a decrease in the number of hindlimb slips in the gridwalk test (Fig. 5G) and a superior recovery in the time it took to sense and then remove a piece of adhesive tape placed on the hindpaws after treatment with CSP–TTK21 (Fig. 5H and fig. S15).
Many of the regenerating axons from the CSP-TTK21–treated mice colocalized with vGlut1, suggesting the formation of putative synapses (Fig. 5, I to K). We then assessed the functionality of these regenerating axons with terminal electrophysiological experiments. We found an increase in the amplitude of compound action potentials recorded above the SCI in treated mice compared to the control group, demonstrating that the Cbp activator increased neural conduction across the lesion site (Fig. 5, L and M). In addition to promoting axon regeneration across the lesion site, we investigated whether CSP-TTK21 also increased sprouting of axons below the level of injury. CSP-TTK21 enhanced the number of vGlut1-positive boutons apposed to motoneurons (putative synapses) in the ventral horn of the lumbar enlargement (Fig. 5, N and O), suggesting spinal circuit reorganization and sprouting of group Ia proprioceptive afferents below the injury. This reorganization of proprioceptive afferents has previously been associated with functional recovery (12, 16). We also observed an increase in the intensity of vGlut1 staining in lamina V of the spinal cord after CSP-TTK21 treatment (fig. S16). In addition, CSP-TTK21 treatment increased H4K8 acetylation within DRG neurons (fig. S17) but did not affect the area or intensity of glial fibrillary acidic protein (GFAP) staining, a marker of the inhibitory astrocytic scar that surrounds the lesion site (fig. S18).

**Pharmacological Cbp/p300 activation enhances sprouting of both descending motor and ascending sensory axons leading to functional recovery after contusion SCI in rats**

To further substantiate the efficacy of CSP-TTK21 in vivo, we evaluated whether CSP-TTK21 could promote anatomical and functional neuroplasticity of motor systems after a more clinically relevant SCI. Adult rats underwent a midthoracic spinal cord contusion (220 kilodynes). CSP-TTK21 or CSP was administered intraperitoneally 6 hours after SCI and repeated weekly thereafter.

To quantify locomotor performance, we applied a principal component (PC) analysis to various parameters (fig. S19) calculated from kinematic recordings...
of quadrupedal walking along a flat corridor. PC1 captured the extent of the recovery, showing that CSP-TTK21 significantly (P = 0.0002) improved locomotor performance compared to CSP-treated rats. Parameters that correlated with improved recovery included reduced paw dragging, increased step height, and more frequent plantar steps with weight bearing (Fig. 6, A to D, and movie S3). The number of footfalls occurring during locomotion across a horizontal ladder also decreased (fig. S20). This functional recovery was associated with increased sprouting of descending reticulospinal and serotoninergic axons within the lumbar spinal cord (Fig. 6, E to J). Consistent with this, CSP-TTK21 enhanced H4K8ac in the reticular formation and raphe nucleus (fig. S21).
CSP-TTK21 also augmented the density of vGlut1-positive boutons from proprioreceptors onto motoneurons located within lumbar segments below the injury (Fig. 6, K and L), which was associated with increased muscle responses evoked by stimulating proprioreceptive afferents (H-reflex; Fig. 6M). However, CSP-TTK21 did not affect the lesion size or Gfap intensity (fig. S22). Together, these results show that activating Cbp using a small molecule promotes sprouting of descending pathways and proprioreceptive afferents below injury and improved recovery of both sensory and motor functions after a contusion SCI in rats.
DISCUSSION

Our work suggests that increasing the neuronal activity of proprioceptive DRG neurons before an injury using EE or chemogenetics elicits Cbp-mediated histone acetylation that is required for an enduring increase in axonal regeneration potential. Activating Cbp using a small molecule at clinically relevant time points in both mouse and rat models of SCI mimicked the effect of increasing neuronal activity. CSP-TTK21 treatment promoted regeneration of ascending sensory axons and sprouting of both sensory and supraspinal motor axons below the lesion. The induced spinal circuit reorganization resulted in electrophysiological and behavioral recovery. The complete lack of EE-mediated increase in neurite outgrowth observed in DRGs from Egr3−/− mice with defective muscle spindle receptors demonstrates the importance of proprioceptive neurons and muscle spindle afferent feedback in triggering the activity-dependent increase in regeneration potential. This finding provides evidence and expands upon the recent demonstration that muscle spindle feedback is essential for inducing the correct anatomical reorganization of projection neurons and functional recovery after SCI (12). In addition to the Egr3−/− mice, we used CTB tracing and PV-tdTomato mice to provide further evidence to support this remarkable cell type specificity. Furthermore, we found that EE drives the expression of genes underlying neuronal activity, calcium signaling, and regenerative pathways in LDNs. The robust gene expression response was largely lost when RNAseq was performed from the whole DRGs that contains multiple neuronal populations and glial cells. This highlights the specificity of EE-mediated gene-expression in LDNs, which mainly constitute proprioceptors and mechanoreceptors. The impact of neuronal activity on axon regeneration could be reproduced experimentally. We showed that the manipulation of DRG neuronal activity using chemogenetics reproduced or abolished the EE-mediated increase in axonal regeneration potential. Furthermore, the increase in DRG neuronal activity alone triggered an increase in axon regeneration, further expanding upon what has been observed recently (38, 39). Together, these data suggest that the effects of EE are essentially elicited by proprioceptive feedback signals, which lead to an enhanced activity of proprioceptive DRG neurons that promotes a lasting augmentation of their regenerative potential. Next, our results provided evidence that the lasting increase in regeneration potential is mediated by a Cbp-dependent increase in histone acetylation and a marked increase in gene expression, including pathways involved in neuronal activity, axonal projection, and cytoskeleton remodeling. Specifically, neuronal activity elicited by EE activates Cbp and increases the acetylation of H4K8. This enduring increase in histone acetylation likely mediates the long-lasting enhancement in regenerative potential of these DRG neurons that extends for several weeks. The levels of histone acetylation are likely maintained because they do not rely solely on the histone deacetylase (HDAC)/acetyltransferase equilibrium but on the overall epigenetic configuration of the locus. Although this is important for histones, it is unlikely for proteins, such as Cbp, whose acetylation status directly depends on the activity of signal transduction pathways. The requirement of Cbp was confirmed by deletion, which completely abrogated the EE-mediated increase in DRG neurite outgrowth and H4K8 acetylation. Furthermore, pharmacological activation of Cbp after SCI promoted axonal regeneration and functional recovery. These data suggest that Cbp is necessary for the EE-mediated increase in DRG neurite outgrowth and that its activation promotes functionally relevant axon regeneration and sprouting, leading to recovery.

Our findings expand upon recent studies by us and others showing that histone acetylation is associated with a transcriptional-dependent enhancement of the regeneration program in neurons (40–45). We previously demonstrated that a conditioning injury activates p300/Cbp-associated factor (Pcaf). Together with HAT p300, Pcaf promotes acetylation of the promoters of known RAGs, which facilitate their expression and thereby enhance axon regeneration after injury (40, 42, 43). The inhibition of HDACs promotes histone acetylation and axonal regeneration (41, 44). Similarly, nuclear export of HDAC5 has been shown to be required for peripheral axon regeneration and for the induction of a number of RAGs (45). Our present results show that EE-dependent histone acetylation does not involve Pcaf because H3K9 acetylation is not altered by EE and does not require p300 because Cbp deletion completely blocked EE-dependent DRG regenerative growth. These observations suggest that a conditioning injury and EE operate via separate signaling mechanisms leading to distinct histone acetylation changes. However, a limitation of the present and previous studies is the lack of systematic screening for posttranslational histone modifications that affect the histone code and gene transcription. The systematic measurement of histone acetylation and methylation could lead to the identification of additional histone-modifying enzymes that modulate EE-dependent or conditioning-dependent axonal regeneration in addition to Cbp/p300 and Pcaf. Collectively, these studies demonstrate the importance of the chromatin environment for the regenerative capacity of DRG neurons. Identifying and manipulating key histone modifiers that can orchestrate broad changes in gene transcription may lead to significant improvements in axon regeneration. The identification of the mechanisms underlying the activity-dependent increase in DRG regenerative growth allowed us to reproduce these effects pharmacologically. We show that the activation of Cbp within a clinically relevant time frame after SCI (within 6 hours) using a nontoxic small molecule promotes regeneration of ascending and descending axons. Cbp activation also triggered a robust sprouting of proprioceptive fibers below the injury, within the lumbar motor circuitry. These changes were associated with enhanced electrophysiological and behavioral functional recovery in sensory and motor tests. Although the specific contribution from each reorganized system remains unclear, we surmise that the reorganization of proprioceptive afferent feedback circuits below the injury is more important to improve precision walking than the relatively short-distance regeneration of ascending fibers. Although the specific mechanisms of recovery require to be studied further, these combined findings show that activity-dependent regenerative pathways triggered preceding an SCI can also be successfully targeted to enhance axon sprouting, regeneration, and sensorimotor recovery after injury.

Rehabilitation strategies including exercises that increase afferent activity in the spinal cord are now well established to augment functional recovery in rodents after SCI, although their effect on axon regeneration in the CNS is not clear (46–48). Moreover, modulation of proprioceptive afferent circuits with electrical stimulation augments neuroplasticity and recovery after an SCI (15, 16, 49, 50). Thus, our results reemphasized the critical role of proprioceptive neurons in steering the reorganization of neural pathways that support functional recovery after SCI.

However, few studies have systematically investigated the impact of task-specific rehabilitation strategies before SCI on
neuroplasticity and functional recovery. One study demonstrated that voluntary exercise before peripheral nerve injury enhances peripheral nerve regeneration (21). This observation is consistent with the robust axon regeneration resulting from an exposure to EE before a peripheral nerve injury. The study by Molteni et al. (21) reported an increase in neurotrophin mRNA in the DRG after exercise. Consequently, they used a pharmacological inhibitor of tropomyosin receptor kinase (Trk) tyrosine kinase to demonstrate that the exercise-mediated increase in DRG outgrowth was contingent on neurotrophin release. These findings differ from our data because we did not observe any significant increase in neurotrophin mRNA or protein levels in DRGs after exposure to EE. Along the same lines, a recent study has shown that exercise after SCI does not change neurotrophin expression in LDNs (51). It is possible that the pharmacological inhibitor of Trk tyrosine kinase may have off-target effects altering multiple intracellular signaling pathways that lead to the reduction in DRG outgrowth. Voluntary exercise has been demonstrated to prevent the reduction of key signaling molecules that are involved in neuroplasticity including pSynapsin I, pCreb, and pCaMK in the spinal cord and brain (52). Similarly, we found that EE triggers an increase in calcium-related signaling molecules known to be important in gene regulation and inducing neuroplasticity, such as Calbindin2 (53). Involuntary exercise was recently shown to promote axon regeneration of propriospinal neurons but not of sensory DRGs after a complete transection of the spinal cord and peripheral nerve graft (54). This suggests that, unlike the Cbp activator used in the present study, increasing the activity of DRG neurons after an SCI using an exercise paradigm may be insufficient to increase the intrinsic regenerative state of the DRGs and promote sensory axon regeneration. In addition, although we did not combine the Cbp activator with neurorehabilitation paradigms, this is worth investigating in the future because they might synergize for improved axonal plasticity and functional recovery. Although we show efficacy of the small-molecule activator of Cbp in two different species and two clinically relevant models of SCI, a limitation is that rodent models of SCI cannot fully replicate the human pathology. In addition, although we did not observe any obvious side effects after treatment with the Cbp activator, these were not systematically assessed. Therefore, future studies will be required to fully evaluate the toxicity profile of the compound before translation to clinical SCI.

Last, it is worth speculating upon the anecdotal evidence that individuals who had an “active lifestyle” recover to a greater degree after SCI than individuals who lived “less active” lifestyles. In addition to the global benefits associated with a healthy lifestyle, our combined observations prompt us to suggest that neurons are “primed” for axonal regeneration and sprouting, which contribute to this enhanced recovery. It will be useful to collect epidemiological data supporting or refuting this hypothesis.

In summary, we have demonstrated an epigenetic-based mechanism underlying activity-dependent neuronal plasticity. The exploitation of this mechanism allowed us to use a pharmacotherapy that enhanced spinal cord repair and functional recovery after SCI, opening a realistic pathway for clinical evaluations.

**MATERIALS AND METHODS**

**Study design**

We investigated the impact of EE on proprioceptive DRG neurons, hypothesizing that EE would prime these neurons to initiate a regenerative response to a subsequent injury. All surgical and experimental procedures on rodents were carried out in accordance with the U.K. Animals (Scientific Procedures) Act 1986 and approved by the veterinarian and ethical committee of Imperial College and the canton of Vaud and Geneva. Animals were assigned randomly to experimental groups, and surgeries were carried out in a random block design. All analyses were performed by the same experimenter who was blinded to the experimental groups. All behavioral testing and analyses were performed by an observer blinded to the experimental groups. All behaviors were replicated two or three times per time point, depending on the experiment. Raw data are in data file S3.

**Statistical analysis**

Results are expressed as mean values ± SEM, and n values represent the number of animals in the experiment. Statistical analysis was carried out using GraphPad Prism 7 (GraphPad, Prism software). The Kolmogorov-Smirnov and Levene’s tests were used to test for normality and the equality of variances. A two-tailed unpaired Student’s t test, one-way ANOVA for evaluation of experiments with more than two groups and one- or two-way repeated-measures ANOVA for functional assessments were used. Tukey’s, Sidak’s, or Fisher’s LSD post hoc tests were applied when appropriate. Behavioral assays were replicated two or three times, depending on the experiment, and averaged per animal. Statistics were then performed over the mean of animals. A threshold level of significance α was set at P < 0.05. Significance levels were defined as follows: *P < 0.05, **P < 0.01, and ***P < 0.001.

**SUPPLEMENTARY MATERIALS**

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Fig. S1. Exposure to EE enhances neurite outgrowth on inhibitory myelin substrate to a similar extent as a conditioning SNA injury.

Fig. S2. Inhibiting transcription with actinomycin-D blocks the EE-mediated increase in DRG neurite outgrowth.

Fig. S3. Exposure to EE enhances axon elongation rather than branching.

Fig. S4. The full EE increases DRG neurite outgrowth compared to the running wheel alone.

Fig. S5. Exposure to EE enhances muscle reinnervation by proprioceptive DRG neurons.

Fig. S6. EE promotes axon regeneration and the formation of putative synapses.

Fig. S7. Neurotrophin and cytokine levels in the DRG and blood serum are not affected by EE.

Fig. S8. RNAseq and proteomic datasets demonstrate that EE strongly modulates pathways involved in neuronal activity, calcium signaling, gene expression, and cytoskeletal changes.

Fig. S9. Efficient transduction and DREADD expression in PV-positive DRGs.

Fig. S10. EE-mediated increase in DRG neurite outgrowth is mediated by neuronal activity.

Fig. S11. H3K4me2, H3K9ac, and H3K4me3 do not change compared to SH.

Fig. S12. Levels of H4K8ac but not of acCbp or pCreb remain elevated in PV-positive DRGs for 5 weeks after exposure to EE.

Fig. S13. Increasing neuronal activity augments the level of H4K8ac and acCbp in DRG neurons.

Fig. S14. The CaMKIIa promoter is active in DRG neurons and drives strong expression of tdTomato after tamoxifen treatment.

Fig. S15. CSP-TTK21 treatment significantly enhances the time to remove adhesive tape placed on the hindpaw.

Fig. S16. CSP-TTK21 treatment promotes sprouting ofafferent fibers below the level of injury.

Fig. S17. CSP-TTK21 treatment enhances H4K8ac in the DRG.

Fig. S18. CSP-TTK21 treatment does not affect the glial scar after a thoracic dorsal spinal cord hemisection in mice.

Fig. S19. List compiling the 78 parameters used for quantifying gait features.

Fig. S20. CSP-TTK21 reduced the number of slips during locomotion along a horizontal ladder.

Fig. S21. CSP-TTK21 treatment enhances levels of H4K8ac in the raphe nucleus and reticular formation.

Fig. S22. CSP-TTK21 treatment does not affect the glial scar after a thoracic contusion SCI in rats.

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Cbp-dependent histone acetylation mediates axon regeneration induced by environmental enrichment in rodent spinal cord injury models


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An epigenetic mechanism for regenerating axons

Functional recovery after spinal cord injury (SCI) is limited by lack of axon regeneration in the mature nervous system. However, recent data showed that increasing neuronal activity promoted axonal regeneration after SCI in rodents. In a new study, Hutson et al. investigated the mechanisms mediating activity-dependent neuronal response in rodent models of spinal cord injury. Increasing neuronal activity using chemical or behavioral approaches promoted recovery through Creb-binding protein (Cbp)–mediated histone acetylation, and using a small-molecule Cbp activator mimicked the effects of increasing neuronal activity. This epigenetic mechanism might be exploited for enhancing repair and functional recovery after SCI.