Spinal Cord Injury in Rats Disrupts the Circadian System

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Significance Statement

Spinal cord injury (SCI) can cause physiologic dysfunction throughout the body. Internal physiologic function is typically synchronized with the environment through the circadian system. Despite the crucial roles of the spinal cord and the circadian system in optimizing whole-body function, it remains unclear whether SCI alters diurnal rhythms. Here, we hypothesized that SCI would disrupt key circadian output and feedback mechanisms. Moderate SCI in rats caused widespread disruption of diurnal measures, including glucocorticoids, body temperature, locomotor activity, and intraspinal clock and inflammatory gene expression. We identify the circadian system as a novel potential target for SCI therapies.

 SCI alters several peripheral oscillators and clocks

Visual Abstract
Spinal cord injury (SCI) perturbs many physiological systems. The circadian system helps maintain homeostasis throughout the body by synchronizing physiological and behavioral functions to predictable daily events. Whether disruption of these coordinated daily rhythms contributes to SCI-associated pathology remains understudied. Here, we hypothesized that SCI in rats would dysregulate several prominent circadian outputs including glucocorticoids, core temperature, activity, neuroinflammation, and circadian gene networks. Female and male Sprague Dawley rats were subjected to clinically relevant thoracic 9 moderate contusion SCI (or laminectomy sham surgery). Diurnal measures—including rhythms of plasma corticosterone (CORT), body temperature, and activity (using small implanted transmitters), and intraspinal circadian and inflammatory gene expression—we were studied prior to and after surgery. SCI caused overall increases and disrupted rhythms of the major rodent glucocorticoid, CORT. Presurgery and sham rats displayed expected rhythms in body temperature and activity, whereas rats with SCI had blunted daily rhythms in body temperature and activity. In parallel, SCI disrupted intraspinal rhythms of circadian clock gene expression. Circadian clock genes can act as transcriptional regulators of inflammatory pathways. Indeed, SCI rats also showed dysregulated rhythms in inflammatory gene expression in both the epicenter and distal spinal cord. Our data show that moderate SCI in rats causes wide-ranging diurnal rhythm dysfunction, which is severe at acute time points and gradually recovers over time. Normalizing post-SCI diurnal rhythms could enhance the recovery of homeostasis and quality of life.

Key words: biological clock; circadian rhythms; diurnal rhythms; neurotrauma; spinal cord injury

Introduction

Spinal cord injury (SCI) dysregulates a constellation of physiologic processes throughout the body. For instance, SCI suppresses peripheral immunity, which increases susceptibility to infection (Schwab et al., 2014). SCI predisposes to accumulating excess adipose tissue and to metabolic syndrome (Jones et al., 2003; Cragg et al., 2015; Sauerbeck et al., 2015). In addition, SCI causes sympathetic dysfunction, which contributes to cardiovascular issues such as autonomic dysreflexia and orthostatic hypotension (Alan et al., 2010; Inskip et al., 2012). Thus, SCI shifts prominent body systems away from a healthy equilibrium (i.e., physiologic homeostasis).

Physiologic homeostasis is regulated by the circadian system, which synchronizes organisms to daily fluctuations in their external environment by temporally organizing internal systems (Fonken and Nelson, 2014). Circadian rhythms are endogenous free-running rhythms (persisting even in constant darkness) lasting ~24 h that are entrainable to external cues; when synchronized by light to day–night cycles, these rhythms are called “diurnal” rhythms. Initial circadian input occurs via light activation of retinal projections to the suprachiasmatic nucleus (SCN). SCN neurons entrain diurnal rhythms in other cells of the brain and body through neural (autonomic), humoral (e.g., glucocorticoids), and physiologic cues (e.g., feeding, activity, and body temperature; Bedrosian et al., 2016). Circadian disruption can impair physiologic function, predispose to disease, and exacerbate postsurgery outcomes (Fonken et al., 2010; Li et al., 2011; Stevens et al., 2014). For example, disrupting the circadian clock with aberrantly timed light exposure following traumatic brain injury in rats increased neuronal death and cortical lesion volume, resulting in worsened sensorimotor and cognitive deficits (Li et al., 2016).

Research in rodents and humans suggests that SCI likely disrupts the circadian system. In mice, SCI transiently increased and disrupted daily expression rhythms of the major rodent glucocorticoid, corticosterone (CORT; Lucin et al., 2007). In rats, thoracic 3 (T3) spinal transection disrupted temperature rhythms for ~14 d post-SCI (West et al., 2015). Together, these results imply that SCI may disrupt diurnal rhythms; however, these studies omitted sham surgery groups, which are essential given that surgery itself can disrupt these systems. Further, these studies did not explore other key circadian outputs (e.g., activity, circadian clock genes). In humans, cervical SCI disrupted diurnal body temperature rhythms at chronic times (Thijssen et al., 2011), and individuals with SCI often experience sleep disturbances (Giannoccaro et al., 2013); yet, diurnal rhythms in humans at acute-to-subacute post-SCI times remain understudied. Moreover, the existence and post-SCI regulation of a molecular circadian clock in spinal cord remains unstudied. Therefore, there is a need to understand the timing and extent of SCI-elicited circadian disruption, and how this could relate to the loss, and potential reacquisition, of physiologic homeostasis.
Here, we hypothesized that SCI in rats would disrupt physiologic, behavioral, and molecular diurnal rhythms. Female and male rats with thoracic SCI of moderate severity displayed a strong increase in, and arrhythmia of, a major circadian entraining factor, CORT. In addition, SCI dampened daily rhythms in body temperature and activity. Moreover, spinal cords from uninjured rats displayed robust diurnal regulation of clock genes; SCI dysregulated intraspinal expression of clock and inflammatory genes, both in epicenter and in distal lumbar spinal cord. We reveal that clinically relevant SCI in rats broadly disrupts circadian function, particularly acutely postinjury.

Materials and Methods

Surgery and animal care
These experiments were approved by University of Colorado Boulder Institutional Animal Care and Use Committee and conformed with ARRIVE (Animal Research: Reporting In Vivo Experiments) standards. In agreement with National Institutes of Health guidelines, male and female rats were included in all experiments. Rats had standard chow and filtered tap water available ad libitum and were maintained on a 12 h light/dark cycle. All surgeries occurred between zeitgeber time 2 (ZT2) and ZT11. Sprague Dawley sham/SCI rats (females, 200–250 × g; males, 320–380 × g; 2–3 months old; Envigo) received isoflurane anesthesia and a T8 laminectomy; SCI rats additionally received a moderate contusion injury (midline SCI; 150 kdyn, 1 s dwell; Infinite Horizon, Precision Systems and Instrumentation). Initial related studies involved pain testing, and inflammation was also studied, so analgesics were not given to any rats for consistency and for limiting confounds (Detloff et al., 2008; Cho et al., 2009; Ellis et al., 2016; Grace et al., 2016). All rats received prophylactic and postsurgery intraperitoneal antibiotics (gentamicin sulfate), subcutaneous saline for 5 d postinjury and a major circadian entraining factor, CORT. In addition, SCI dampened daily rhythms in body temperature and activity. Moreover, spinal cords from uninjured rats displayed robust diurnal regulation of clock genes; SCI dysregulated intraspinal expression of clock and inflammatory genes, both in epicenter and in distal lumbar spinal cord. We reveal that clinically relevant SCI in rats broadly disrupts circadian function, particularly acutely postinjury.

Tissue processing, histology, and analysis
To evaluate lesion size and spared tissue area, tissue was collected for immunohistochemistry (n = 7 male and female rats) at 7 dpi. Rats received an intraperitoneal pentobarbital overdose and were transcardially perfused with 0.9% saline, then 4% paraformaldehyde. Spinal cords were suspended in paraformaldehyde overnight, cryoprotected in 36% sucrose, and cryosectioned (16 μm; Gaudet et al., 2015). For immunohistochemistry, slides were incubated with 10% normal donkey serum (1 h), then with primary antibodies (overnight; mouse anti-glial fibrillary acidic protein (GFAP; 1:100; catalog #0869110, MP-Biomedicals) and rabbit anti-Iba1 (1:1000; catalog #019-19741, Wako Chemicals), then with secondary antibodies (2 h; Alexa Fluor-488 donkey anti-mouse (A-21202) and Alexa Fluor-546 donkey anti-rabbit (A-10040); both 1:500; Thermo Fisher Scientific and DAPI (nuclear stain; catalog #D1306, Thermo Fisher Scientific). Images were captured on an Olympus IX81 Microscope and analyzed using Fiji (Schindelin et al., 2012).

Locomotor testing
Locomotor recovery was assessed using the Basso, Beattie, and Bresnahan (BBB) scale (Basso et al., 1995; female sham: n = 6; SCI, n = 5; male sham: n = 7; SCI, n = 5) by two condition-blind observers before surgery; and at 1, 4, 7, 10, 14, 21, 28, 35, and 42 d postsurgery.

Collection and measurement of plasma corticosterone
Rats were pair housed; cage mates had the same surgery (at start: female sham: n = 6; SCI, n = 10; male sham: n = 6; SCI, n = 10; two female and one male SCI rat died). Rats were acclimated to handling for 5 d; then, blood samples were collected from immobilized unanesthetized rats via tail nick (<500 μL/24 h). Blood was collected presurgery, and 2, 7, and 14 d postsurgery at ZT0, ZT6, ZT12, and ZT18 (starting at ZT0; CORT was typically the lowest). For dark-phase collections, dim red light headlamps were used (Fonken and Nelson, 2014). Samples were centrifuged (10,000 × g; 10 min) to isolate plasma. Plasma was used in a CORT ELISA (catalog #ADI-901-097, Enzo). Several samples (various rats and time points) had insufficient plasma, so were excluded, and samples that took longer than 3 min to collect were excluded (due to confounding stress-elicited CORT).

Body temperature and locomotor activity measurement
Individually housed rats were implanted intraperitoneally with radiotelemetric transmitters (MiniMitter, Respiration; Fonken et al., 2012). After a 1 week recovery, prelaminectomy baseline activity/temperature was recorded for 1 week. Home cages on 12 TR-4000 receiver boards linked with DP-24 DataPorts (MiniMitter) continuously collected temperature/activity. Activity and body temperature were recorded for >6 weeks postsurgery.

Female and male experiments were completed separately due to a limited number of telemetry receivers. First, males [sham, n = 6; SCI, n = 5 (originally 6); one SCI rat transmitter did not transmit] were tested. Including shams was important: basic surgeries can disturb circadian rhythms (Farr et al., 1988). Sham/SCI rats received daily handling postsurgery (twice-daily bladder emptying or handling for shams; at approximately ZT2 and ZT10) and daily antibiotic/saline injections for 5 d postsurgery (approximately ZT2). Two weeks postsurgery, rats voided bladders independently, and daily checks continued without handling. After the male experiment, the experimental design was improved. For females (sham, n = 6; SCI, n = 6), presurgery handling (twice per day) acclimated the rats and reduced potential stress effects on postsurgery activity/temperature. Post surgery antibiotic/saline injections (for 5 d postsurgery) occurred at afternoon (ZT10) animal care to limit the disruption of key early inactive phase rhythms.

Diurnal regulation of gene expression
For the uninjured PCR study, tissues were collected from uninjured female/male rats across the day (ZT0, ZT6, ZT12, and ZT18; n = 6/sex/time point). Rats received an intraperitoneal pentobarbital overdose then were per-
fused with 0.9% saline. Tissues were flash frozen for PCR (L4–L5 was used for uninjured spinal cord analyses; T8 (lesion) and L4–L5 were used for postsurgery analyses). L4–L5 was used as a distal spinal cord site because it integrates hindpaw nociceptive information, which could provide useful information for future studies. L4–L5 was used for the uninjured analyses, since the T8 injury site was expected to have major shifts in gene expression. The lumbar spinal cord was predicted to have more minor SCI-elicited changes. For the sham/SCI PCR study, female/male rats received sham/SCI surgery (lesion) and L4–L5 were used for postsurgery analyses. (L4–L5 was used for uninjured spinal cord analyses; T8 spinal cord contusion (150 kdyn, 1 s dwell) integrates hindpaw nociceptive information, which could provide useful information for future studies. L4–L5 was used for the uninjured analyses, since the T8 injury site was expected to have major shifts in gene expression. The lumbar spinal cord was predicted to have more minor SCI-elicited changes. For the sham/SCI PCR study, female/male rats received sham/SCI surgery (lesion) and L4–L5 were used for postsurgery analyses. (L4–L5 was used for uninjured spinal cord analyses; T8 spinal cord contusion (150 kdyn, 1 s dwell).

Statistics

Data were analyzed (SigmaPlot 13.0, SYSTAT) using Student’s t test or nonparametric Mann–Whitney U test; or ANOVAs (one-, two-, or three-way ANOVA, as appropriate). Holm–Sidak post hoc tests were completed when appropriate for all tests involving more than two groups. Circadian data were analyzed using Circ Wave ([https://www.euclock.org/results/item/circ-wave.html](https://www.euclock.org/results/item/circ-wave.html)), which assesses independent circadian data (i.e., one individual contributes to a single data point) to identify waveforms and acrophases (rhythm peaks). Twice-daily postsurgery handling (voiding/injections) caused stress-elicited hyperthermia and hyperactivity in all groups and were excluded from Circ Wave analyses for clarity. Activity data were also analyzed using ClockLab 6.0 (Actimetrics); actograms and wavelets were assessed for each rat, and data from representative male sham and SCI individuals are presented. Actograms were scaled between 0 and 1. Data were plotted as the mean ± SEM.

Results

T8 spinal cord contusion (150 kdyn, 1 s dwell) causes tissue pathology and locomotor deficits

First, we examined tissue damage in male and female rats at 7 dpi in this SCI model (Gaudet et al., 2017; Fig. 1).
As expected, substantial tissue loss was observed at the T8 lesion epicenter (Fig. 1D); moving rostrocaudally away from the epicenter, there was progressively more tissue sparing (Fig. 1E). There were no significant differences in lesion size or tissue sparing in females versus males (lesion volume: females, 8.15 ± 0.70 mm³; males, 7.48 ± 0.46 mm³; p > 0.05). Locomotor recovery after sham/SCI was assessed (Fig. 1F). At 1 dpi, average BBB scores indicated that SCI rats had movement of one hindlimb joint. By 42 dpi, SCI rats recovered frequent hindlimb stepping with no (or little) coordination (females: BBB score, 10.6 ± 0.6; males: BBB score, 11.0 ± 1.4).

Spinal cord injury causes a transient increase and arrhythmia in plasma CORT

CORT release is regulated in a circadian manner; this steroid hormone maintains metabolic homeostasis across the day and helps to entrain extra-SCN circadian rhythms (Nicolaides et al., 2014). Here, we assessed whether SCI dysregulated plasma CORT levels and rhythms (Fig. 2). Before surgery, female and male rats showed peak trough patterns in plasma CORT (Fig. 2A; D’Agostino et al., 1982): CORT had cycle peak (acrophase) near early to mid-active phase (at approximately ZT12; females, ZT12: 1.4 ± 0.3 ng/ml; males, ZT12: 0.6 ± 0.1 ng/ml), and was lowest at the start of the inactive phase (ZT0; females: 0.6 ± 0.2 ng/ml; males: 0.4 ± 0.1 ng/ml). CORT levels were higher overall in presurgery females than males (Cavigelli et al., 2005). In females, CORT was unusually high at ZT6, which could have been related to a handling-elicited CORT increase (Gärtner et al., 1980). CORT was expressed rhythmically before injury in both females (F(2,37) = 3.465, p < 0.05) and males (F(2,33) = 6.647, p < 0.005).

SCI disrupted typical rhythms and significantly increased CORT levels at acute postinjury times (females: main effect of injury: F(3,62) = 3.606, p < 0.05; males: main effect of injury: F(3,70) = 22.584, p < 0.001). Females with SCI had increased CORT at 7 dpi (vs presurgery; at ZT0; p < 0.05); males with SCI showed significant CORT increases at 2 dpi (vs presurgery; at ZT0 and ZT6; p < 0.001) and 7 dpi (at ZT0; p < 0.05; Fig. 2B). For average CORT concentration across the day, SCI in female and male rats caused significant increases in average CORT at 2 d postsurgery (females: vs presurgery, p < 0.05; males: main effect of surgery; also at 2 dpi vs both presurgery and sham, p < 0.001; Fig. 2C).

Spinal cord injury alters daily rhythms in body temperature

Body temperature and activity are two prominent outputs of the circadian system that also help to entrain...
circadian rhythms in cells throughout the body (Reebs and Mrosovsky, 1989; Buhr et al., 2010). To measure these parameters, rats were implanted with a small transmitter and were studied before sham/SCI, and from acute-to-chronic times postsurgery. (The studies on male and female rats were completed consecutively, due to the limited number of telemetry receivers; for details, see Materials and Methods.)

Before surgery, female and male rats displayed the expected diurnal rhythms in body temperature: core temperature was higher during the active phase, and lower during the inactive phase (female-SCI presurgery acrophase, ZT16.6; male-SCI presurgery acrophase, ZT17.3; Fig. 3). Immediately after surgery (i.e., after replacing postsurgery rats on transmitter boards), SCI but not sham rats experienced hypothermia (lasted ~18 h; ANOVA main effect: female: $F_{(1,155)} = 31.664, p < 0.001$; male: $F_{(1,142)} = 24.663, p < 0.001$; and significant differences at specific times; Fig. 3).

After initial post-SCI hypothermia, female rats showed significant SCI-elicited temperature dysregulation. By 2 d postsurgery, female sham and SCI rats both showed approximately typical rhythms in core temperature (Circ Wave; all showed rhythmicity). Compared with sham rats, female SCI rats had shifted (advanced) acrophases between 2 and 6 d postsurgery and increased inactive phase temperatures between 2 and 5 d postsurgery (2–6 dpi SCI acrophases: ZT12.0, ZT12.9, ZT13.0, and ZT14.5, respectively; ANOVA of main effect of treatment and injury–time interaction, 2–3 d postsurgery: $F_{(1,203)} = 6.678, p < 0.05$; sham vs SCI, 2 d postsurgery: $F_{(1,155)} = 31.664, p < 0.001$; male: $F_{(1,142)} = 24.663, p < 0.001$; and significant differences at specific times; Fig. 3).

Figure 2. SCI in female and male rats disrupts diurnal rhythms in CORT. A, Before surgery, female and male rats exhibited diurnal rhythms in CORT, with nadir levels at the beginning of the inactive (light) phase and peak levels toward the beginning of the active (dark) phase. B, SCI increased CORT levels and altered CORT rhythms at 2 and 7 dpi; CORT rhythms were normalized by 14 dpi. C, Average CORT level across the day in female and male SCI rats was increased at 2 dpi. *ZT differences or SCI vs presurgery, ANOVA with Holm–Sidak post hoc test, $p < 0.05$; †SCI vs sham at that time point, $p < 0.05$. New Research 6 of 18

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Figure 3. SCI in female and male rats disrupted diurnal rhythms of core body temperature. Before surgery, female and male rats exhibited typical diurnal core temperature rhythms. Immediately after surgery, SCI rats displayed hypothermia. SCI disrupted diurnal
VT3, VT6, and VT9; 3 d postsurgery: VT3, VT6, and VT9, $p < 0.05$; 4–5 d postsurgery: $F_{(1,203)} = 12.207, p < 0.01$; sham vs SCI: 6 d postsurgery: VT18, $p < 0.05$; inactive phase temperature: SCI > sham, 15 d postsurgery, $p < 0.05$. SCI rats displayed no significant differences versus sham at 7, 13, or 14 d postsurgery (7 dpi: ANOVA, no main effect of treatment, $F_{(1,203)} = 0.982$; 13–14 dpi: ANOVA, no main effect of treatment, $F_{(1,203)} = 1.147, p > 0.05$).

Male rats showed a similar pattern to females of SCI-elicited temperature dysregulation, but with somewhat delayed recovery of rhythms. Sham rats at 2–3 dpi showed approximately typical rhythms in core temperature (2 and 3 d postsurgery sham acrophases: VT13.3 and VT16.2; all showed rhythmicity); conversely, male SCI rats at 2–3 dpi did not have significant core temperature rhythms (ANOVA main effect of treatment: $F_{(1,186)} = 0.688, p > 0.05$). Male SCI rats displayed shifted acrophases and increased inactive phase temperatures compared with sham rats between 4 and 5 d postsurgery (acrophases: sham, 4 d postsurgery: 16.4; SCI, 4 dpi: 13.7; sham, 5 d postsurgery: 16.4; SCI, 5 dpi: 12.5; ANOVA main effect of treatment: $F_{(1,186)} = 0.657, p > 0.05$; inactive phase temperature: SCI > sham, 4–5 d postsurgery, $p < 0.05$). There were no significant differences caused by SCI in core temperature at 6–7 d postsurgery (ANOVA, no main effect of treatment: $F_{(1,186)} = 0.400, p > 0.05$) nor at 13–14 d postsurgery (ANOVA, no main effect of treatment: $F_{(1,186)} = 2.794, p > 0.05$).

To understand more broadly whether SCI influenced temperature rhythms, daily temperature was processed in ClockLab in grouped multiday “bins” (Fig. 4). Before surgery, the peak daily temperature phase occurred in females at an angle of 263.5° (or VT17.8) and in males at 275.2° (or VT18.3). After surgery, females with SCI showed no significant difference in phase from shams ($p > 0.05$, two-way repeated-measures ANOVA with Holm–Sidak post hoc test). In contrast, males with SCI displayed significantly advanced temperature phases: SCI males had earlier temperature acrophase starting at 2–5 d postsurgery [ANOVA main effect of treatment: $F_{(1,81)} = 43.299, p < 0.001$; 2–5 d postsurgery: male sham rats, phase angle 261.9° ± 3.4° (VT17.5); male SCI rats, phase angle 208.3° ± 15.3° (VT13.9); $p < 0.05$], and these SCI advanced rhythms continued through all 5 d bins examined through 26–30 d postsurgery [26–30 d postsurgery: male sham rats, phase angle 268.7 ± 4.9° (VT17.9); male SCI rats, phase angle 249.9° ± 4.1° (VT16.7); $p < 0.05$]. At 31–35 d postsurgery and beyond, male SCI rat temperature acrophases were not significantly different from those of male sham rats [e.g., 31–35 d postsurgery: male sham rats, 269.5° ± 3.5° (VT18.0); male SCI rats, 257.2° ± 4.0° (VT17.1); $p > 0.05$].

Overall, SCI altered the average core temperature at early postinjury times—particularly due to increased temperature during the inactive phase—and showed substantial recovery at subacute times. SCI males (vs females) displayed significant and longer-lasting shifts in temperature acrophase.

Spinal cord injury disrupts amount and diurnal rhythms of activity

Activity patterns are a strong index of the integrity of the circadian system. Although SCI clearly reduces activity, it is also possible that it disrupts activity rhythms. Rhythmic activity is important for entraining peripheral circadian rhythms (Reebs and Mrosovsky, 1989; Tahara et al., 2017). Before surgery, rats showed typical diurnal patterns of activity, with more activity during the active phase (female SCI rats presurgery acrophase, VT15.1; male SCI rats presurgery acrophase, VT17.9; counts in active phase: females, 65 ± 2%; males, 67 ± 2%).

![Temperature peak: phase angle](image-url)

**Figure 4.** SCI significantly advanced temperature acrophase in males, but not females. Temperature phase angles (in bins of 4–5 d) were studied in female and male rats before surgery, then up to 50 d postsurgery. **A.** Compared with sham rats, females with SCI showed no significant difference in phase angle after surgery. **B.** SCI males had significantly advanced temperature acrophases from 2–5 d through 26–30 d postsurgery. **C.** Phase plot showing how SCI affected the phase angle over time postsurgery (up to 25 d for clarity). Angle indicates the time of acrophase; radial distance depicts time postsurgery. *$p < 0.05$.*

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**continued**

rhythms in core temperature between 2 and 5 d postsurgery. Diurnal temperature regulation was similar between sham and SCI rats by 13–14 d postsurgery. *$p < 0.05$ for sham vs SCI at that time point, ANOVA with Holm–Sidak post hoc test.*
After surgery, differences in activity between sham and SCI rats occurred mainly during the active phase (Fig. 5). Sham rats showed relatively typical activity levels and diurnal activity rhythms by 2–3 d postsurgery (2 and 3 d postsurgery sham acrophases: female, ZT15.2 and ZT16.2; male, ZT15.8 and ZT16.8). In contrast, female and male SCI rats had strongly reduced activity throughout the day at 2 dpi (total counts: female sham rats, 13,000 ± 2000; female SCI rats, 7000 ± 2000; p < 0.05; male sham rats, 13,200 ± 800; male SCI rats, 3200 ± 800; p < 0.001). Although this pattern of post-SCI reduced activity continued for days, there was some recovery of diurnal activity rhythms over the first 5 dpi. At 6–7 dpi, female and male rats with SCI showed further recovery of activity, but not 6 dpi, although their overall activity remained significantly lower than that of sham rats. Patterns of activity at 6 d postsurgery and beyond were not significantly different between sham and SCI rats in both females and males (percentage counts in active phase at 6 d postsurgery: female sham rats, 70 ± 2%; female SCI rats, 63 ± 3%; p > 0.05; male sham rats, 70 ± 2%; male SCI rats, 62 ± 3%; p > 0.05).

At 13–14 d postsurgery, female and male SCI rats had activity patterns that were more similar to shams (acrophases: female sham rats, 13.9; female SCI rats, 13.3; male sham rats, 16.2; male SCI rats, 15.8; total counts in active phase at 14 dpi: female sham rats, 9000 ± 900; female SCI rats, 7000 ± 1000; p > 0.05; male sham rats, 8500 ± 500; male SCI rats, 6600 ± 500; p < 0.05). These patterns were even more similar at a chronic post-SCI time point (acrophases: female sham rats, 15.8; female SCI rats, 15.7; male sham rats, 16.1; male SCI rats, 15.5). Therefore, SCI transiently dampens activity levels, and the timing of post-SCI recovery of daily activity counts and rhythms parallels the recovery of other circadian measures.

Spinal cord injury delays recovery of 24 h activity rhythms

To establish postsurgery latency to recovery of 24 h activity rhythms, actogram and wavelet analyses were performed. These qualitative assessments help visualize diurnal rhythms. Data from one male sham rat and one male SCI rat (both representative) are presented. Before surgery, the rats displayed expected rhythms in activity (Fig. 6A). After sham surgery, rats recovered relatively typical rhythms within ~1 d postsurgery. In contrast, after SCI, rats recovered 24 h activity rhythms approximately 6–7 d postsurgery (Fig. 6B, C, yellow arrows). Therefore, SCI substantially delayed the recovery of diurnal activity rhythms.

Spinal cord injury disrupts clock gene expression in epicenter and lumbar spinal cord

Rhythms of circadian “clock” gene expression exist throughout the body; however, clock gene expression has not been systematically characterized in the spinal cord. Thus, first we sought to establish how clock genes are regulated by time of day in the uninjured rat spinal cord (Fig. 7A). Our data reveal that clock genes are expressed rhythmically in the L4–L5 spinal cord. The “core clock” components Per2, Cry1, and Bmal1 were all regulated in spinal cord by time and sex (Cry1, Bmal1: main effect of time (p < 0.001) and sex (females > males, p < 0.001); Per2: significant time–sex interaction, F_{3,44} = 2.953, p < 0.05) and were expressed rhythmically across the day (all in both females and males; acrophases: Per2: female, ZT15.2 ± 1.2; male, ZT12.2 ± 1.2; Cry1: female, ZT16.3 ± 1.4; male, ZT16.8 ± 1.5; Bmal1: female, ZT23.0 ± 1.4; male, ZT0.8 ± 1.3). Similarly, Rev-erbα, a transcription factor in a secondary circadian feedback loop, was regulated in spinal cord by time of day (interaction with sex: F_{3,44} = 3.953, p < 0.05) and was rhythmically expressed (in females and males; Rev-erbα acrophases: female, ZT6.0 ± 1.4; male, ZT6.6 ± 1.1).

To establish whether SCI alters molecular circadian rhythms, rats were subjected to T8 sham (laminectomy) or SCI surgery, and tissue was collected at 2 d postsurgery in the light (ZT6) or dark (ZT18) phase. Clock gene expression was assessed in T8 epicenter and lumbar spinal cord (Fig. 7B,C). In T8 epicenter, SCI (vs sham surgery) substantially downregulated key clock genes Per2, Cry1, Bmal1, and Rev-erbα (F_{1,45} = 45.7, 113.2, 82.1, and 135.6, respectively; all p < 0.001; main effect at ZT6 and ZT18; Fig. 7B). In addition, there were time-of-day differences in sham rats that were abolished by SCI in several genes (Per2, Cry1, and Rev-erbα; post hoc test), and there were sex differences in Cry1 and Rev-erbα expression (main effect; males higher).

Dysregulation of circadian genes by SCI extended beyond the lesion site. Lumbar spinal cord from SCI (vs sham) rats displayed significantly reduced expression of Per2, Cry1, Bmal1, and Rev-erbα (F_{1,30} = 40.7, 48.4, 62.2, and 23.8, respectively; all p < 0.001; main effects; SCI also downregulated Per2 at ZT6 and ZT18; Fig. 7C). In addition, there were main effects on clock gene expression of time (Cry1, Bmal1, and Rev-erbα) and sex (Per2, Cry1, and Bmal1). Additional clock genes dysregulated by SCI included Per1 and Clock (Table 2).

Spinal cord injury disrupts inflammatory gene expression in epicenter and lumbar spinal cord

Inflammatory genes in the CNS can also be regulated by time of day, and this likely confers differential immunocompetence across the day (Fonken et al., 2015). In the healthy adult rat spinal cord, several inflammatory genes were expressed differentially across the day (Fig. 8A): the proinflammatory cytokines IL-1b (main effect of time: F_{3,43} = 3.54, p < 0.05; females had significant daily rhythm, p < 0.05) and Tnfα (main effect of time: F_{3,44} = 5.57, p < 0.005; males had significant daily rhythm, p < 0.01); the inflammatory cytokine IL-6 (main effect of time: F_{1,44} = 6.99, p < 0.001; males had significant daily rhythm, p < 0.005); and the microglial/macrophage activation marker Cd88 (significant interaction of time and sex: F_{3,44} = 5.51, p < 0.005; males had significant daily rhythm, p < 0.005). SCI significantly dysregulated inflammatory gene expression in epicenter and lumbar spinal cord (main effects; Fig. 8B,C). IL-1b was significantly upregulated in SCI (vs sham) rats in the T8 lesion epicenter but was
Figure 5. SCI in female and male rats disrupted diurnal rhythms in activity. Data show cumulative activity counts per 3 h time period. Before surgery, female and male rats exhibited typical diurnal rhythms in activity, with rats more active in the dark phase. As expected,
SCI rats were less active at acute times postinjury (up to 7 d postsurgery). Interestingly, at subacute (14 d postsurgery) to chronic times, SCI rats regained activity levels and diurnal rhythms in activity that were similar to those of sham rats. *p < 0.05 for sham vs SCI at that time point, ANOVA with Holm–Sidak post hoc test.

downregulated in the lumbar spinal cord ($F_{(1,45)} = 11.9$ and $F_{(1,30)} = 144.7$, respectively; both $p < 0.001$; main effects). *Tnfa* was upregulated by SCI in both the epicenter and the lumbar spinal cord ($F_{(1,45)} = 12.9$ and $F_{(1,29)} = 7.9$, respectively; both $p < 0.001$; main effects; also main effect of time: lumbar). *IL-6* expression was increased in SCI rats in the epicenter, but was reduced in the lumbar spinal cord ($F_{(1,44)} = 43.6$, $p < 0.001$; $F_{(1,29)} = 4.5$, $p < 0.05$, respectively; main effects; also main effect of sex in both tissues). *Cd68* expression was increased by SCI in

![Graphs showing diurnal activity rhythms](image)

**Figure 6.** Rats with SCI show delayed recovery of diurnal activity rhythms. These representative data are from one male sham and one male SCI rat. **A, B,** Actograms display continuous measures of activity across the day (ZT; x-axis) and over time postsurgery (y-axis). **A,** Before surgery, these rats in the sham and SCI groups had expected diurnal patterns of activity (i.e., increased activity between ZT12 and ZT24; higher black bars). **B,** After surgery, sham rats quickly recover more typical activity rhythms, whereas SCI rats show delayed postsurgery recovery of rhythms. Yellow arrows highlight approximate latency to the recovery of 24 h rhythms; also shown in **C. C,** Wavelet analysis shows that before surgery rats display strong 24 h rhythms, and that sham and SCI rats have different postsurgery latencies to recover 24 h activity rhythms (yellow arrows). The sham rat recovered 24 h rhythms within ~1 d postsurgery, whereas the SCI rat recovered 24 h rhythms at ~6 d postsurgery. The intensity of rhythm across days is represented by the color continuum: purple (minimal rhythm) through blue and green to red (intense rhythm).
A **Uninjured: spinal cord**

![Image of clock gene expression in uninjured rats.](image)

**Diurnal regulation of clock genes in spinal cord from uninjured rats and from sham/SCI rats at 2 d postsurgery.** A, Female and male rat spinal cords express clock genes, and clock gene expression varies through the day. Per2, Cry1, Bmal1, and Rev-erba displayed rhythmic expression in the spinal cord. Females generally expressed higher levels of expression of clock genes. B, SCI disrupts clock gene expression at the lesion epicenter. Clock gene expression at ZT6 and ZT18 was assessed. Sham spinal cords showed clock gene variation between the two time points (e.g., Per2, Cry1, and Rev-erba). After SCI, injury epicenters showed strongly reduced expression and ablated diurnal variation of these four clock genes, suggesting circadian disruption. C, SCI disrupts clock gene expression in spinal cord distal to injury. The lumbar spinal cord, which was not directly injured by contusion, showed decreased expression of the four clock genes examined in SCI vs sham rats. There were also time-of-day differences for all genes presented, and a significant main effect of sex for Per2, Cry1, and Bmal1. Black — indicates that females or males for that gene show significant rhythm; red + indicates injury difference (sham vs SCI), p < 0.05; yellow hourglass indicates time difference, p < 0.05; blue gender symbol indicates sex difference, p < 0.05. Symbols at the top of each graph indicate the significant main effect; symbols above/below data indicate significant interactions.

![Table 2: Expression of additional clock (Per1, Clock) and inflammatory (Iba1, Mhc II) genes at 2 d postsurgery in the lesion epicenter and lumbar spinal cord (data only for genes not shown in figures).](table)

| Gene   | Female | Male | Significant differences | Interactions |
|--------|--------|------|-------------------------|--------------|
|        | Sham   | SCI  | ZT6 | Sham | SCI | ZT6 | Sham | SCI | ZT18 | Sham | SCI | ZT18 | Sham | SCI | ZT18 |
| Per1   |        |      |     |      |     |     |      |     |      |      |     |      |      |     |      |
| Lesion | 1.00 ± 0.10 0.39 ± 0.06 & 1.19 ± 0.07 0.23 ± 0.03 & 1.01 ± 0.17 0.35 ± 0.06 & 2.01 ± 0.57 0.34 ± 0.01 & SCI; ZT6 vs ZT18 & Sham × ZT |
| Lumbar | 1.00 ± 0.05 1.04 ± 0.29 & 0.97 ± 0.09 0.70 ± 0.03 & 0.78 ± 0.08 0.57 ± 0.07 & 0.84 ± 0.12 0.50 ± 0.02 & SCI, sex & N.S. |
| Clock  |        |      |     |      |     |     |      |     |      |      |     |      |      |     |      |
| Lesion | 1.00 ± 0.16 0.32 ± 0.03 & 1.21 ± 0.18 0.27 ± 0.04 & 0.90 ± 0.18 0.36 ± 0.09 & 1.56 ± 0.33 0.29 ± 0.03 & SCI & Sham × ZT |
| Lumbar | 1.00 ± 0.03 0.71 ± 0.10 & 1.07 ± 0.06 0.64 ± 0.04 & 0.94 ± 0.07 0.53 ± 0.07 & 0.83 ± 0.08 0.45 ± 0.04 & SCI & Sham × ZT |
| Iba1   |        |      |     |      |     |     |      |     |      |      |     |      |      |     |      |
| Lesion | 1.00 ± 0.21 0.75 ± 0.03 & 0.86 ± 0.08 0.93 ± 0.09 & 0.69 ± 0.09 0.41 ± 0.07 & 0.59 ± 0.04 0.78 ± 0.14 & SCI × ZT; ZT6 × inj. |
| Lumbar | 1.00 ± 0.07 1.36 ± 0.04 & 0.96 ± 0.08 1.38 ± 0.13 & 1.32 ± 0.05 2.06 ± 0.45 & 1.23 ± 0.12 1.54 ± 0.18 & N.S. |
| Mhc II |        |      |     |      |     |     |      |     |      |      |     |      |      |     |      |
| Lesion | 1.00 ± 0.17 0.21 ± 0.05 & 0.82 ± 0.09 0.21 ± 0.03 & 0.65 ± 0.14 0.23 ± 0.05 & 0.79 ± 0.14 0.22 ± 0.05 & SCI & N.S. |
| Lumbar | 1.00 ± 0.19 0.50 ± 0.05 & 0.54 ± 0.21 0.54 ± 0.18 & 1.59 ± 0.38 0.86 ± 0.14 & 1.09 ± 0.30 0.54 ± 0.11 & SCI, sex & N.S. |

For statistical comparisons (far right column): SCI, significant main effect of SCI vs sham; ZT, significant main effect of time (ZT6 vs ZT18); sex, significant main effect of sex; Sham × ZT, significant interaction between sham and ZT; N.S., no significant difference.
the epicenter, but was downregulated in the lumbar spinal cord (F(1,44) = 32.6 and F(1,29) = 31.2, respectively; both p < 0.001; main effects; also main effect of time (epicenter and lumbar) and sex (epicenter); SCI also altered intraspinal expression of Iba1, an RNA expressed by microglia/macrophages, and Mhc II, an antigen-presenting molecule expressed by microglia/macrophages; Table 2). Thus, SCI robustly dysregulated intraspinal clock and inflammatory gene expression.

**Discussion**

This study used female and male rats to determine whether SCI dysregulated acute-to-chronic physiologic and behavioral rhythms and related molecular outputs. Moderate T8 spinal cord contusion dysregulated plasma CORT—a key humoral output of and feedback signal for the circadian system—particularly at 2 and 7 d postsurgery. In addition, SCI dampened diurnal rhythms in locomotor activity and body temperature. SCI strongly decreased the expression of several clock genes in the epicenter and lumbar spinal cord, and also altered the expression of intraspinal inflammatory genes at two times of day. Therefore, moderate SCI in a rat model severely disrupts daily rhythms in CORT, activity, body temperature, and intraspinal gene expression. The physiologic parameters studied gradually reacquired more typical daily rhythms over time. Our results suggest that moderate contusion SCI causes widespread, transient disruption of physiologic homeostasis (Fig. 9).

**Locomotor recovery after SCI**

As expected, rats with SCI showed a substantial deficit in hindlimb movement at 1 dpi, and the rats recovered frequent stepping by the chronic 42 dpi time point. Locomotor recovery had not previously been assessed in this model (a T8 150 kdyn contusion with 1 s dwell; Putatunda et al., 2014; Gaudet et al., 2017; Hook et al., 2017). Interestingly, the post-SCI locomotor recovery presented here complements the findings of others: our average BBB score of 10.6 in female 42 dpi rats supports that this injury severity is between the severities reported previ-
SCI disrupts diurnal rhythms and physiologic homeostasis

Diurnal rhythms help to optimize organism function across the day; thus, disrupting diurnal rhythms likely shifts the physiology of an animal away from homeostasis. Prolonged diurnal disruptions have detrimental effects for the individual. Here, we identify SCI as a disruptor of several prominent physiologic rhythms: CORT, core temperature, and activity.

Glucocorticoids are steroid hormones that help to synchronize extra-SCN circadian rhythms (Balsalobre et al., 2000). This is possible because SCN-derived signals entrain CORT rhythms to light and SCN cells lack CORT receptors, whereas most non-SCN cells express CORT receptors. In this manner, CORT can modulate peripheral clocks directly (e.g., CORT–receptor complex drives Per1 and reduces Rev-erbα transcription; Oster et al., 2006; Dickmeis, 2009; Surjit et al., 2011). CORT is potently upregulated by immune activation and helps to resolve inflammatory responses; however, CORT is also critically involved in stress responses (Frank et al., 2013) and basic homeostatic functions (Nicolaides et al., 2014). Given these roles of CORT in health and pathology, we hypothesized that SCI would dysregulate CORT rhythms. Indeed, male and female rats with acute SCI had increased CORT levels and disrupted CORT rhythms. Previous work supports our findings: female mice with T9 contusion SCI had acute, transient dysregulation of glucocorticoid rhythms (Lucin et al., 2007), although no sham mice were included. A recent study by Pruss et al. (2017) found that SCI in mice increased serum CORT levels (T1 SCI > T9 SCI; 3 dpi CORT with no time of day specified); similarly, humans within 96 h post-SCI showed increased serum cortisol levels. SCI in mice and humans substantially increases susceptibility to peripheral infection (pneumonia). In mice, SCI-elicited CORT increases could be quenched by removing the major source of systemic CORT, the adrenal glands, but adrenalectomy did not reduce the frequency of pneumonia (Pruss et al., 2017).

Interestingly, transplanting denervated adrenal glands into adrenalectomized SCI mice normalized CORT levels and limited susceptibility to pneumonia (Pruss et al., 2017; but also see Oster et al., 2006). Thus, accelerating the recovery of CORT rhythms could help to regain homeostasis, including diurnal rhythms.

We also revealed that daily rhythms in body temperature were significantly altered at acute post-SCI times and gradually recovered. Immediate post-SCI hypothermia lasted ~24 h (hypothermia not observed in shams). Body temperature acrophase was particularly disrupted in male rats with SCI, which had altered temperature acrophase that persisted through 30 dpi. Others also observed SCI-elicited temperature arrhythmia in rats (West et al., 2015). Similarly, humans with acute trauma (including SCI) commonly experience hypothermia, which can copresent with other pathologies and lead to death (Kirkpatrick et al., 1999). In humans with chronic SCI, diurnal regulation of core body temperature was disrupted after cervical, but not thoracic, SCI (although acute post-SCI rhythms were not assessed; Thijsen et al., 2011).

Body temperature rhythms help coordinate peripheral circadian rhythms (Buhr et al., 2010); thus, optimizing post-SCI recovery of body temperature rhythms could accelerate the recovery of homeostasis.
Activity rhythms were transiently dampened by SCI, which could have more widespread implications for the circadian system. For example, timed activity can be used to help strengthen diurnal rhythms: in mice with a mutation in core circadian clock machinery, scheduled (late night) wheel running strengthened molecular and physiologic rhythms (Schroeder et al., 2012). Behavioral activity feeds back to the SCN: exercise in mice suppresses SCN neuron activity, and exercising at the proper time of the cycle increases SCN diurnal rhythm amplitude (van Oostervout et al., 2012). In humans, exercising at abnormal times (during the night) phase shifts diurnal rhythms (Youngstedt et al., 2016). Therefore, post-SCI diurnal rhythms may be strengthened by optimizing the time of day of key activities, including rehabilitation, social activity, and feeding.

Glucocorticoids, body temperature, and activity modulate circadian oscillators throughout the body, so SCI-elicited dysregulation could alter daily rhythms more broadly. Indeed, typical diurnal rhythms in body function can be disrupted in humans by SCI (e.g., sleep; Giannoccaro et al., 2013) and by related medical interventions (Li et al., 2011), so expediting the recovery of diurnal rhythms after SCI could improve neuroprotection and functional outcomes.

It is important to note that we had limited telemetry receivers, so we completed the body temperature and activity transmitter study separately in males and females. In addition, limitations were noted after completing the male study and methods were improved for the female portion (e.g., rats were acclimated to twice-daily handling before surgery), which could have affected the consistency of the results. Despite this, the results were remarkably similar between males and females, both for sham and SCI rats, and differences existed between sham and SCI groups. Our experience and data also highlight that typical postsurgery rodent care and behavioral testing in many research laboratories (e.g., twice-daily rodent care during the light/inactive phase) likely further disrupts daily rhythms (as well as sleep), and could be a confound to the factor under study (McEwen, 2006). Future SCI studies could improve their design by having the daily injections and major care immediately before the active phase, and by having a second potential daily care time during the dark phase or at another minimally disruptive time (e.g., using shifted light schedules in animal housing, if helpful). In addition, given that intense rodent care during the beginning of the inactive phase is likely a strong disruptor of rhythms, future studies could compare whether this type of disruption affects anatomic and neurologic outcomes compared with a "circadian-optimized" care schedule. These findings would have translational relevance, since early post-SCI patients in hospitals likely experience excess circadian disruptions (that could be improved with relatively simple interventions).

Another distinction between studies described herein is that the telemetry study was completed with individually housed rats (to enable measuring the temperature and activity of a single rat), whereas all other studies were completed with rats housed in pairs. This is important to note, given that housing rats individually is a stressor (Von Frijtag et al., 2000; Westenbroek et al., 2005) and can impair recovery after CNS injury (Craft et al., 2005; Kareлина et al., 2009; Gaudier-Diaz et al., 2017). Furthermore, single housing may reduce relative activity (vs having a cage mate) and therefore likely worsens other aspects of recovery (e.g., locomotor recovery; Van Meeteren et al., 2003).

**SCI alters intraspinal expression and diurnal regulation of clock and inflammatory genes**

At 2 d postsurgery, SCI remarkably reduced the expression of clock genes observed in the lesion epicenter, and a more modest (but significant) decrease also occurred in the lumbar spinal cord. In the lesion epicenter, this is likely affected by altered cell composition: sham T8 spinal cords contain typical CNS cellular constituents, whereas SCI T8 epicenters include various activated immune, glial, and other cells (Gaudet and Fonken, 2018). Epicenter-localized cells could have different clock gene levels that cause the massive decrease observed. Another possibility is that SCI dysregulates a finely tuned circadian and inflammatory gene expression network: immune cell-specific circadian gene dysregulation can provoke inflammatory responses (Nguyen et al., 2013; Fonken et al., 2016; Segal et al., 2018); conversely, the induction of cytokines such as TNF-α and IL-1β can suppress the expression and/or function of circadian genes (Cavadini et al., 2007). Indeed, intraspinal Tnfα and IL-1β diurnal regulation inversely correlated with Per2 expression, both in uninjured spinal cord and in the SCI epicenter. Similarly, Rev-erbα reduces IL-6 expression (Gibbs et al., 2012); here, diurnal variation in Rev-erbα and IL-6 levels were inversely related in uninjured spinal cord, and the SCI-elicited decrease in epicenter Rev-erbα paralleled an increase in IL-6. These findings underscore the potential relevance of clock and inflammatory gene cross talk. In addition, SCI downregulated clock genes even distal to lesion in lumbar spinal cord, where cell infiltration likely has limited influence. Overall, decreased and arrhythmic clock gene expression by SCI could broadly regulate immune and homeostatic function in the CNS.

Inflammatory gene expression was differentially affected by SCI in epicenter and lumbar spinal cord. In the epicenter, most genes assessed were upregulated; conversely, most genes assessed in the lumbar spinal cord were downregulated. L4–L5 spinal cord was chosen as a distal spinal cord site because it integrates hindpaw nociceptive information. At later times post-SCI, excess below-level intraspinal inflammation likely contributes to SCI-elicited neuropathic pain (Hains and Waxman, 2006; Detloff et al., 2008), and we found that this contusion model causes neuropathic pain symptoms by 14 dpi in female and male rats (Gaudet et al., 2017). The reduced expression of inflammatory markers at 2 dpi may represent an initial remote intraspinal dampening of the immune response, which may precede chronic neuroinflammation that exacerbates neuropathic pain and pathology.
Here, two time points [middle of the light (ZT6) and dark (ZT18) phase] were used to assess clock and inflammatory gene expression at 2 d post-SCI. This is a limitation of the study, since with two time points it is not possible to make broader conclusions about the effects of SCI on circadian phase, amplitude, and average expression. Future studies could examine the expression at additional times of day. Further, the disruption of circadian outputs persisted beyond 2 dpi, including rhythms of serum CORT, body temperature, and activity (all were disrupted until at least 7 dpi), yet our study only assessed gene expression changes at 2 d postsurgery. Gene expression was assessed at 2 d postsurgery because robust changes in the circadian system were seen at this time, and because early changes in lesion pathology and inflammation likely have strong effects on recovery trajectory (Noble et al., 2002; Kigerl et al., 2009; Gaudet et al., 2018; McCready et al., 2018). It is also important, however, to understand how clock and inflammatory gene patterns shift through subacute and chronic times postsurgery. In the future, studying circadian gene expression changes at additional times post-SCI would provide more information about the robustness and duration of the effects of SCI on the circadian system.

**Future directions**

Our study highlights that SCI disrupts whole-body functions, yet unresolved questions remain about post-SCI circadian disruption and related potential therapies. As shown in Figure 9, SCI alters the regulation of several circadian outputs. Daily rhythms are entrained by light input to the retina, influencing the SCN, which is likely protected from changes caused by SCI [e.g., unlike most cells of the body, SCN neurons lack CORT receptors; Balsalobre et al., 2000; Woodruff et al., 2016]. However, peripheral rhythms are synchronized and entrained by other circadian oscillators, including CORT, body temperature, activity, and autonomic circuits (Buijs et al., 2016), that are perturbed by SCI. Therefore, the disruption of circadian outputs could contribute to more widespread circadian disruption throughout the body. Here, a moderate T8 spinal cord contusion was used; future studies could establish whether more severe injuries or injuries at other spinal levels (e.g., cervical or high thoracic, which would more strongly perturb autonomic function; Inskip et al., 2009) differentially alter postinjury circadian dynamics. In addition, our study did not address whether circadian disruption was simply a result of injury, or whether disruption itself further exacerbated outcomes. Future studies could determine whether amplifying post-SCI rhythms using, for example, timed exercise, feeding, light/dark schedule manipulations, and injections of circadian-related drugs, could improve recovery measures (discussed more below). The strengthening of post-SCI rhythms could also influence other long-term recovery measures, such as survival, neuroprotection, chronic pain, autonomic function, and general health. Further, since altering circadian rhythms affects other aspects of metabolism, metabolic cage telemetry and calorimetry (Gaudet et al., 2016) could reveal whether boosting post-SCI rhythms improves short-term and long-term metabolic health. Finally, to better understand the effect of SCI on circadian rhythms, SCI researchers could adopt other models and strategies frequently used in circadian research, such as the *PER::luciferase* transgenic rodent, which uses the clock gene promoter from *Per* to express the bioluminescent *luciferase* gene and enables studying rhythm synchrony/dysregulation *in vitro* (Yamazaki et al., 2000) and *in vivo* (Tahara et al., 2012).

If SCI-elicited circadian disruption worsens recovery and metabolism, amplifying diurnal rhythms using entraining strategies soon after injury could expedite recovery (Roenneberg and Merrow, 2016). For instance, improving the darkness of rooms during nights (e.g., minimize brightness of lights/monitors; use red lights and filters) and incorporating bright morning light (e.g., having an outdoor-facing window in the room; Engwall et al., 2014), feeding during the day (Stokkan et al., 2001; Fonken et al., 2010), and optimizing the time of day for rehabilitation/activity (Schroeder et al., 2012) could improve sleep–wake cycles and re-entrain diurnal rhythms. Using these and other strategies to accelerate post-SCI recovery of homeostasis could boost key early recovery processes and overall postinjury outcomes.

**Conclusions**

In conclusion, we used a clinically relevant rat spinal contusion model to assess how SCI affects circadian dynamics. We established that moderate thoracic SCI has broad effects on diurnal rhythms, including disrupted rhythms of plasma CORT levels, activity, body temperature, and intraspinal gene expression. SCI caused robust diurnal rhythm disruption at acute post-SCI times (2 and 7 dpi; across various circadian measures); SCI rats recovered more typical diurnal rhythmicity by subacute-to-chronic times (14–42 dpi). SCI-associated disruption of these key regulators of physiologic homeostasis may feed back to impede SCI recovery. Future discovery and clinical SCI studies could incorporate measures of circadian function, which may reveal post-SCI “chronotherapies” that help to regain homeostasis and improve recovery.

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