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Reorganization of spinal neural connectivity following recovery after thoracic spinal cord injury: insights from computational modelling

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eLife Assessment

This **important** study describes a computational model of the rat spinal locomotor circuits and how they could be plastically reconfigured after lateral hemisection or contusion injuries to replicate gaits observed experimentally in vivo. Overall, the simulation results **convincingly** mirror the gait parameters observed experimentally. The model suggests the emergence of detour circuits after lateral hemisection, whereas after a midline contusion, the model suggests plasticity of left-right and sensory inputs below the injury.

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Abstract

Rats exhibit significant recovery of locomotor function following incomplete spinal cord injuries, albeit with altered gait expression and reduced speed and stepping frequency. These changes likely result from and give insight into the reorganization within spared and injured spinal circuitry. Previously, we developed computational models of the mouse spinal locomotor circuitry controlling speed-dependent gait expression (Danner et al. 2017; Zhang et al. 2022). Here, we adapted these models to the rat and used the adapted model to explore potential circuit-level changes underlying altered gait expression observed after recovery from two different thoracic spinal cord injuries (lateral hemisection and contusion) that have roughly comparable levels of locomotor recovery (Danner et al., 2023). The model reproduced experimentally observed gait expression before injury and after recovery from lateral hemisection and contusion, and suggests two distinct, injury-specific routes to restored function. First, recovery after lateral hemisection required substantial functional restoration of damaged descending drive and long propriospinal connections, suggesting compensatory plasticity through formation of detour pathways. Second, recovery after a moderate midline contusion predominantly relied on reorganization of spared sublesional networks and altered control of supraspinal cervical circuits, compensating for weakened propriospinal and descending pathways. Despite these differences, sensitivity analysis revealed that restored activation of sublesional lumbar rhythm-generating circuits and appropriately balanced lumbar commissural connectivity are the key determinants of post-injury gait expression, suggesting that injury symmetry shapes how the cord reorganizes, but effective recovery in both cases depends on re-engaging these lumbar networks, which makes them prime targets for therapeutic intervention.

Introduction

Although incomplete spinal cord injuries partially disrupt communication across the lesion, they are often followed by substantial recovery of locomotor function (Burns and Ditunno, 2001; Kuerzi et al., 2010). These improvements are accompanied by reorganization of spared and severed descending pathways and spinal circuitry (Raineteau and Schwab, 2001; Fouad and Tse, 2008; Takeoka et al., 2014; Filli and Schwab, 2015; Engmann et al., 2020; Lemieux et al., 2024). However, it remains unclear which plastic changes of neural connectivity underlie functional recovery and if the symmetry of the lesion influences those changes.

Locomotion is primarily controlled by spinal circuitry (Grillner, 1981; Orlovsky et al., 1999; Kiehn, 2006, 2016; McCrea and Rybak, 2008; Rybak et al., 2015), which produces the locomotor rhythm and pattern, and controls interlimb coordination in response to supraspinal inputs. Flexor-extensor alternation in each limb is regulated by rhythm generators located bilaterally within the lumbar and cervical enlargements (Kato, 1990; Ballion et al., 2001; Juvén et al., 2012; Rybak et al., 2015; Danner et al., 2016, 2017; Frigon, 2017; Fig. 1A). These rhythm generators receive descending inputs from the brainstem, which modulate locomotor frequency and speed (Danner et al., 2016; Capelli et al., 2017; Caggiano et al., 2018; Josset et al., 2018; Ausborn et al., 2019), and are interconnected by commissural (left–right) and long propriospinal (lumbar–cervical) neuronal pathways, which define interlimb coordination (Talpalar et al., 2013; Danner et al., 2016, 2017; Ruder et al., 2016). Thoracic spinal cord injuries disrupt these descending pathways and inter-enlargement long propriospinal neuronal connections, while leaving the sublesional lumbar and supraspinal cervical circuitry largely intact (Fig. 1B,C). Thus, the animals’ ability for speed-dependent gait expression following post-injury recovery can provide insights into the state of reorganization of descending and inter-enlargement pathways (Shepard et al., 2021, 2023; Danner et al., 2023).

Indeed, following thoracic lateral hemisection or moderate midline contusion injuries, rats recover weight-bearing overground locomotion, though with a reduced maximal speed and altered speed-dependent gait patterns (Danner et al., 2023). Specifically, post-hemisection recovery allows expression of a subset of pre-injury gaits — walk, trot, canter, and gallop — in a speed-dependent manner, albeit with reliance on the contralesional limb as lead in non-alternating gaits. In contrast, mild-to-moderate midline contusion injuries result in a more pronounced speed reduction, loss of non-alternating gaits, and emergence of novel alternating gait patterns (Danner et al., 2023).

We developed a series of computational models of the spinal circuitry that controls speed-dependent interlimb coordination and gait expression (Danner et al., 2016, 2017; Ausborn et al., 2019, 2021; Zhang et al., 2022). By reproducing the speed-dependent gait patterns of intact mice (Bellardita and Kiehn, 2015; Lemieux et al., 2016) and simulating the effects of silencing or ablating various classes of commissural (Talpalar et al., 2013; Bellardita and Kiehn, 2015) and long propriospinal interneurons (Ruder et al., 2016; Zhang et al., 2022), these models offer insights into the organization and function of spinal locomotor circuitry, including the roles of commissural and long propriospinal interneurons as well as the broader interactions between brainstem drive and the spinal circuitry that regulate gait expression. Given the similarity in gait patterns between rats (Danner et al., 2023) and mice (Bellardita and Kiehn, 2015; Lemieux et al., 2016), we adapted these models to rats as a foundation for studying post-injury plasticity.

Here, we use a computational model of spinal circuitry to investigate potential reorganization of neural connectivity underlying locomotor control following recovery from symmetrical thoracic contusion and asymmetrical (lateral) hemisection injuries. The model reproduces speed-dependent gait expression in rats before injury and after recovery from hemisection and contusion. In our model, reproduction of altered gait expression observed after recovery from the fully asymmetrical hemisection injury required substantial functional restoration of severed long propriospinal connections and descending drive. This suggests the involvement of compensatory mechanisms, such as local axon sprouting and rewiring, resulting in formation of detour pathways

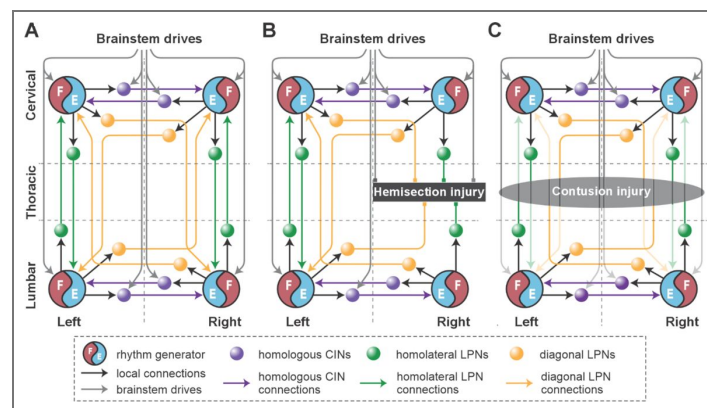


Figure 1. Model concept.

A. Organization of the spinal locomotor circuitry (intact). B, C. Neural structures affected by thoracic hemisection (B) or contusion (C) injuries. Spheres represent neural populations involved in commissural and long propriospinal pathways. Descending drives and synaptic interactions are shown by arrowheads. Decreased color intensity in C signifies partial disruption of pathways by contusion injury. CINs, commissural interneurons; LPNs long propriospinal interneurons.

via the intact hemicord. In contrast, after a symmetrical contusion injury, the model suggests that the observed post-recovery gait changes rely on strengthened intra-enlargement connectivity, increased afferent input, and/or changes in intrinsic excitability of the sublesional circuitry that compensate for weakened long propriospinal connections and descending drive.

Results

Model of the rat spinal locomotor circuitry

The intact (pre-injury) model (Figs 1A and 2) was based on our previous models of mouse speed-dependent gait expression (Danner et al., 2017; Zhang et al., 2022) and was adapted to reproduce rat locomotion. Like the previous models, it consists of four rhythm generators (RGs), located on the left and right sides of the cervical and lumbar compartments (left and right fore and left and right hind RGs). Each RG includes a flexor and extensor center (F and E) that mutually inhibit each other via inhibitory interneuron populations (InF and InE). Both flexor and extensor-centers incorporate a persistent (slowly-inactivating) sodium current (I_{NaP} , see Methods), allowing them to intrinsically generate rhythmic activity under certain conditions defined by the level of excitation, which is regulated by descending brainstem drive. The flexor centers operate in a bursting mode, while the extensor centers receive relatively high drive that keeps them in the mode of tonic activity and exhibit rhythmic activity only due to rhythmic inhibition from the corresponding flexor centers. Thus, each RG produced alternating flexor-extensor activity defining two major locomotor phases (flexor and extensor). The frequency of RG oscillations was defined by brainstem drive to each flexor center; it increased with increasing drive, primarily due to shortening of the extensor phase durations. This reflects the asymmetric modulation of stance and swing phase durations observed in limbed locomotion across species (Halbertsma, 1983; Hildebrand, 1989; Bertram, 2016), including rats (Cohen and Gans, 1975; Danner et al., 2023), where increases in locomotor frequency are primarily achieved by shortening stance (extensor) phases while swing (flexor) phases remain relatively stable — a pattern also generated intrinsically by spinal locomotor circuits across species (Frigon and Gossard, 2012; Danner et al., 2015; Shevtsova et al., 2015).

The left and right RGs within the cervical and lumbar compartments interact via a series of homologous commissural pathways mediated by commissural ($V0_D$, $V0_V$, and $V3$) and local ($V2a$, Ini and $Ini1$) interneurons, coordinating the activities of the left and right homologous RGs. The cervical and lumbar RGs interact via a series of homolateral ($Sh2$ and $LPNi$) and diagonal ($dV0_D$, $dV0_V$ and $aV3$) long propriospinal pathways, coordinating the fore-hind RG activities. Homologous (intra-enlargement) $V0_D$ and $V0_V$ and descending diagonal $V0_D$ interneurons receive the inhibitory brainstem drive, allowing them to coordinate left–right and fore–hind activities in a speed-dependent manner. The organization of spinal locomotor circuitry was based on multiple sets of experimental findings on speed-dependent gait expression of intact mice and after manipulating various classes of local, commissural, and long propriospinal interneurons (Talpalár et al., 2013; Bellardita and Kiehn, 2015; Rybak et al., 2015; Shevtsova et al., 2015; Danner et al., 2016, 2017, 2019; Ruder et al., 2016; Ausborn et al., 2021; Zhang et al., 2022). The detailed description of the proposed connectome of spinal circuitry can be found in Danner et al. (2017) and Zhang et al. (2022). The connection weights and neuron parameters in the current model were adjusted to reproduce characteristics of locomotor behavior in rats (see Methods).

We quantified frequency-dependent changes in inter-RG coordination and flexor/extensor phase durations in response to variations in brainstem drive (parameter α , see Methods) by calculating normalized phase differences between activities of pairs of RGs and compared the resulting gait expression to rat locomotion. Specifically, inter-RG coordination in the model was compared to interlimb coordination in rats, flexor and extensor phase durations were compared to swing and stance phase durations, and the period of oscillation was compared to the step-cycle duration in rats. These parameters were used to calculate gait in both the model and in rats to assess frequency-dependent gait expression (see Methods).

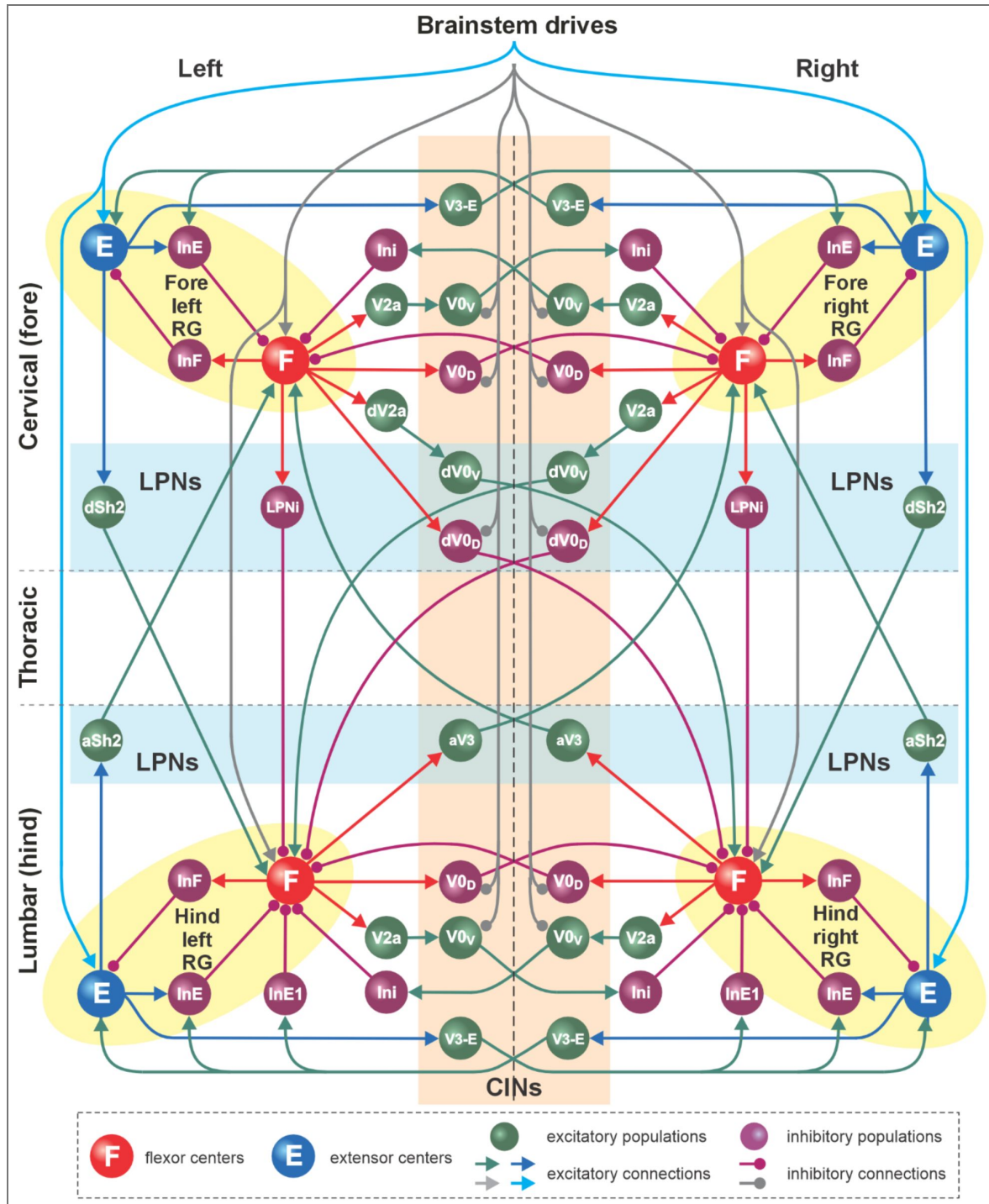


Figure 2. Detailed model schematic.

Spheres represent neural populations. Excitatory drives and connections are marked by arrowheads; inhibitory connections are marked by circles. RG, rhythm generator; In, interneuron; CINs, commissural interneurons; LPNs, long propriospinal neurons; a-, ascending; d-, descending. See text for details. Adapted from Danner et al. (2017) and Zhang et al. (2022) with modifications.

Model reproduces gait expression of intact rats

To ensure suitability of the model to study potential circuit reorganization following recovery from spinal cord injury, we first compared speed-dependent gait expression of the model in the intact case (intact or pre-injury model) with that of intact rats while traversing a 3-m long tank (Danner et al., 2023). Locomotor bouts consisting of periods of acceleration and deceleration were simulated by successively and repetitively ramping the brainstem drive (parameter α , see Methods) from a value at which trot is usually expressed ($\alpha=0.55$) to one at which bound is expressed ($\alpha=1.05$) and back down; step-to-step variability was insured by including a noisy current ($\sigma_{\text{Noise}} = 1.1$ pA; see Methods).

Representative examples of bouts of locomotor activities of the intact model and rat are shown in Fig. 3A1 and 3A2, respectively. Step frequencies increased progressively in both the model and rats, driven primarily by shortening of the extensor and stance phases (Fig. 3A1 and 3A2, upper panels). As frequency increased, the hind left–right normalized phase differences in the model and rats (orange lines in Fig. 3A1 and 3A2, bottom panels) gradually shifted from alternation (~ 0.5 phase difference) to synchronization ($\sim 0/1$ phase difference). Meanwhile, homolateral phase differences remained close to alternation (green), and diagonal phase differences transitioned from synchronization to alternation (blue). As a result, both the model and rat transitioned from trot to gallop and then half-bound gallop with increasing frequency. As shown previously (Danner et al., 2023), speed-dependent gait changes were gradual, rather than abrupt, resulting in the expression of transitional gaits (Fig. 3A).

The average gaits identified in the intact model (Fig. 3B1) — trot, canter, transverse and half-bound gallops (with left and right lead), and bound — closely matched those observed in rats (Fig. 3B2). In the model, trot was characterized by synchronized diagonal RG activity and alternating left–right activation of homologues fore and hind RGs. Canter exhibited synchronization of only one pair of diagonal RGs. Gallops showed quasi-synchronization of the hind RGs and a non-zero phase difference between the fore RGs. Bound was characterized by left–right synchronization of the fore and hind RGs and alternation of the homolateral and diagonal RGs. Note the striking similarity of average extensor/stance phases and normalized phase differences in modeling and experimental results (Fig. 3B1 and 3B2).

To compare frequency-dependent gait expression between the model and rats across bouts, normalized phase differences between pairs of RGs (Fig. 4A1) and corresponding pairs of limbs (Fig. 4A2) were plotted against locomotor frequency. In both cases, the frequency-dependent gait transitions from trot through gallop to half-bound gallop and bound (Fig. 4B1, B2) can be seen as progressive changes in the normalized phase differences (Fig. 4A1, A2). At low frequencies (~ 3 – 4 Hz), left–right alternation of the hind RGs/hindlimbs and the fore RGs/forelimbs dominated. At medium frequencies (~ 4 – 5.5 Hz), left–right phase differences showed progressive change from alternation to synchronization in both directions up to essential synchronization, while some steps still maintained left–right alternation. Finally, at the highest frequencies (>6.5 Hz in model and >6 Hz in rats), most steps exhibited left–right fore and hind phase differences close to 0 or 1 (synchronized activity). The normalized homolateral phase differences remained alternating, close to 0.5 across the frequency range. The normalized diagonal phase differences gradually changed from synchronized to alternating. In parallel, phase durations showed asymmetric frequency-dependent shortening, with a more pronounced reduction in extensor (stance) phase duration relative to flexor (swing) phase duration, consistent between the model (Fig. 4C1) and rats (Fig. 4C2).

Trot in the intact model was expressed at oscillation frequencies from about 3 to 6 Hz (Fig. 4B1) and was the most prevalent gait in our simulations ($\sim 45\%$, Fig. 5A1), which is similar to the intact rats (Figs 4B2 and 5A2). The transverse gallop, both in the model and in rats, was expressed at slightly higher frequencies (~ 4.5 to 6.5 Hz) and accounted for $\sim 30\%$ of all steps, making it the second most prevalent gait (Figs 4B1, B2 and 5A1, A2). Half-bound gallop was observed in the model at ~ 5 to ~ 7 Hz (Fig. 4B1), similar to rats (Fig. 4B2), and was observed in $\sim 10\%$ of all steps (Fig. 5A1, A2). Bound was expressed at ~ 5 to 7 Hz in both the model and rats

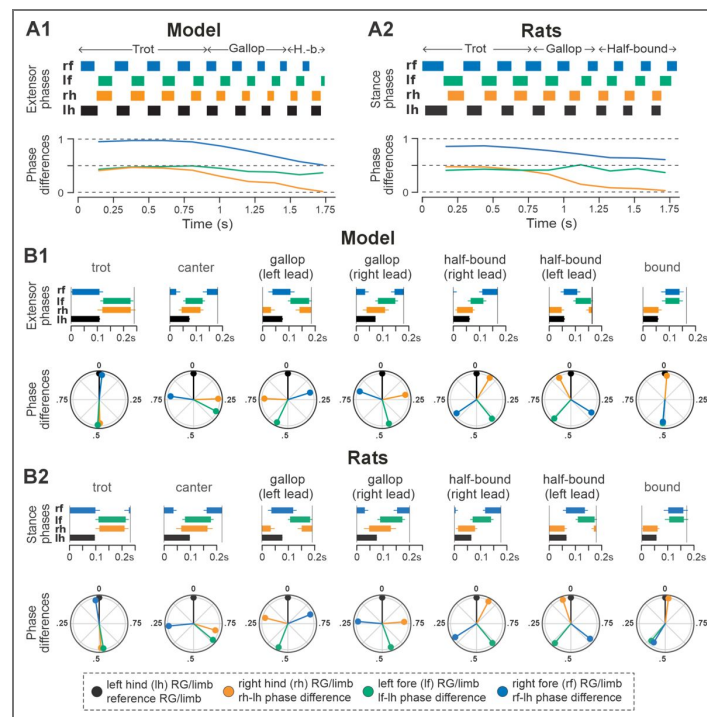


Figure 3. Gait expression in pre-injury/intact model and rats.

A1, A2 Extensor/stance phases (upper panels) and instantaneous normalized phase differences (bottom panels) of representative bouts for the model (A1) and a rat (A2). B1, B2 Average extensor/stance phases for each gait (upper panels) and circular plots of average normalized phase differences for each gait (bottom panels) expressed in the intact model (B1) and rats (B2). Detailed statistical results for rats are reported in [Danner et al. \(2023\)](#). H.-b., Half-bound; RG, rhythm generator.

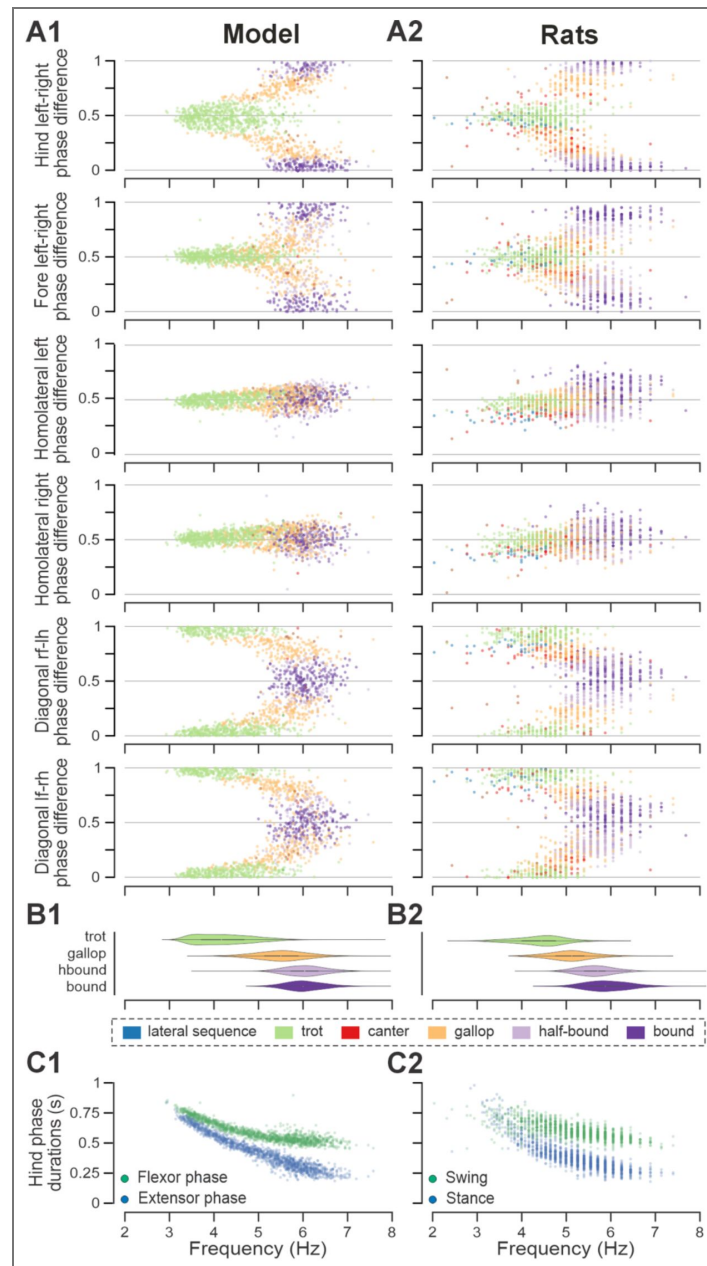


Figure 4. Frequency-dependent distribution of normalized phase differences, gaits, and phase durations in the model and in rats.

A1, A2 Scatter plots of normalized phase differences against frequency of locomotor oscillations in the intact model (A1) and rats (A2). Each dot represents one step cycle. Gaits are classified for each step cycle and color-coded. B1, B2 Distribution of gaits vs. locomotor frequency in the model (B1) and rats (B2). Due to the low prevalence of lateral-sequence and canter steps, these gaits were omitted in B1 and B2. C1, C2 Flexor and extensor phase duration in the model (C1) and duration of swing and stance in rats (C2) against frequency of locomotor oscillations. Same number of step cycles are shown for the model and animals; model step cycles were randomly sampled. l-, left; r-, right; -f, fore RG/limb; -h, hind RG/limb; RG, rhythm generator.

and accounted for ~10% of steps (Figs 4B1,B2 and 5A1,A2). Canter was the least prevalent gait in the intact model (~0.5% of steps; not shown in Fig. 4B1), and also occurred rarely, mostly as a transitional gait, in rats (Danner et al., 2023). Although the exact proportions of the expressed gaits in the intact model and rats differ (Fig. 5A1,A2), their frequency-dependent distribution is qualitatively similar (compare Fig. 4B1 and 4B2). Importantly, both in the model and intact rats, the ranges of locomotor frequency where particular gaits were expressed overlapped (Fig. 4B1,B2) and transitions between gaits occurred most prevalently between gaits in the sequence of their frequency-dependent expression (Figs 4B1 and 5C1,D1), similar to the gait transitions in the intact rats (Fig. 4B2 and Fig. 2 in Danner et al., 2023).

Finally, to assess the step-to-step variability while accounting for variance introduced by gait changes, we measured the mean deviations from the circular exponential moving averages of each phase difference per bout (Danner et al. 2023). Both the model and rats exhibited a similar pattern: the mean deviations of the moving averages of the two left-right and diagonal phase differences were comparable to each other and consistently higher than those of the two homolateral phase differences (Fig. 5B1,B2).

In summary, the model reproduces the experimental results on locomotion of intact rats well: both the model and rats show progressive gait transitions with frequency increase (from trot to gallop and then to half-bound gallop and to bound) on a continuum in phase-space, and gaits were expressed in overlapping frequency ranges.

Modelling recovery from lateral thoracic hemisection injury

Following recovery from the asymmetrical lateral thoracic hemisection, rats regain the ability to locomote over a wide range of speeds but lose the ability to perform the fastest gaits (half-bound gallop and bound) and exhibit clear left-right asymmetry in interlimb coordination, predominantly using the contralesional limb as the lead during asymmetric gaits such as canter and gallop (Danner et al., 2023). Nevertheless, the expressed gaits represent a subset of intact gaits with appropriate interlimb coordination. Here, we used the model to investigate potential reorganization of spinal locomotor circuitry that could underlie the observed speed-dependent gait expression following recovery from hemisection.

To do this, we first modeled the acute effect of the injury by eliminating severed connections from the model (Figs 1B and 6A). Specifically, right thoracic hemisection was modeled by eliminating brainstem drive to the ipsilesional lumbar compartment and all long propriospinal neuron (LPN) connections on the side of the injury. Note that axons of contralaterally projecting LPNs generally cross the midline at a spinal level close to the cell body (Reed et al., 2006; Pocratsky et al., 2020), thus we assumed that contralaterally projecting LPNs with contralesional cell bodies and homolaterally projecting LPNs with ipsilesional cell bodies are axotomized by the thoracic hemisection.

Then, by manually tuning model parameters, we sought to identify a small set of connectivity changes sufficient to reproduce speed-dependent gait expression observed experimentally in rats following recovery from hemisection injury (Fig. 6B). Specifically, we assumed that the injury-affected LPN and descending pathways, along with local cervical and lumbar circuitry, undergo plasticity associated with functional recovery, e.g., through regrowth (sprouting), detour pathways, synapse loss/formation, or upregulation of sensory feedback, and that descending control of lumbar and cervical circuitry through uninjured pathways is altered.

These assumptions were implemented in the model (Fig. 6B) by:

- i. (functional) recovery of the injury-affected LPN connections to 40% of their pre-injury weights,
- ii. recovery of drive to the ipsilesional hind rhythm generator to 90% of its pre-injury strength (this might be partially due to concomitant upregulation of sensory input),
- iii. recovery of inhibitory drive to the ipsilesional lumbar V0_v commissural interneuron (CIN) to 50% of its pre-injury strength, and

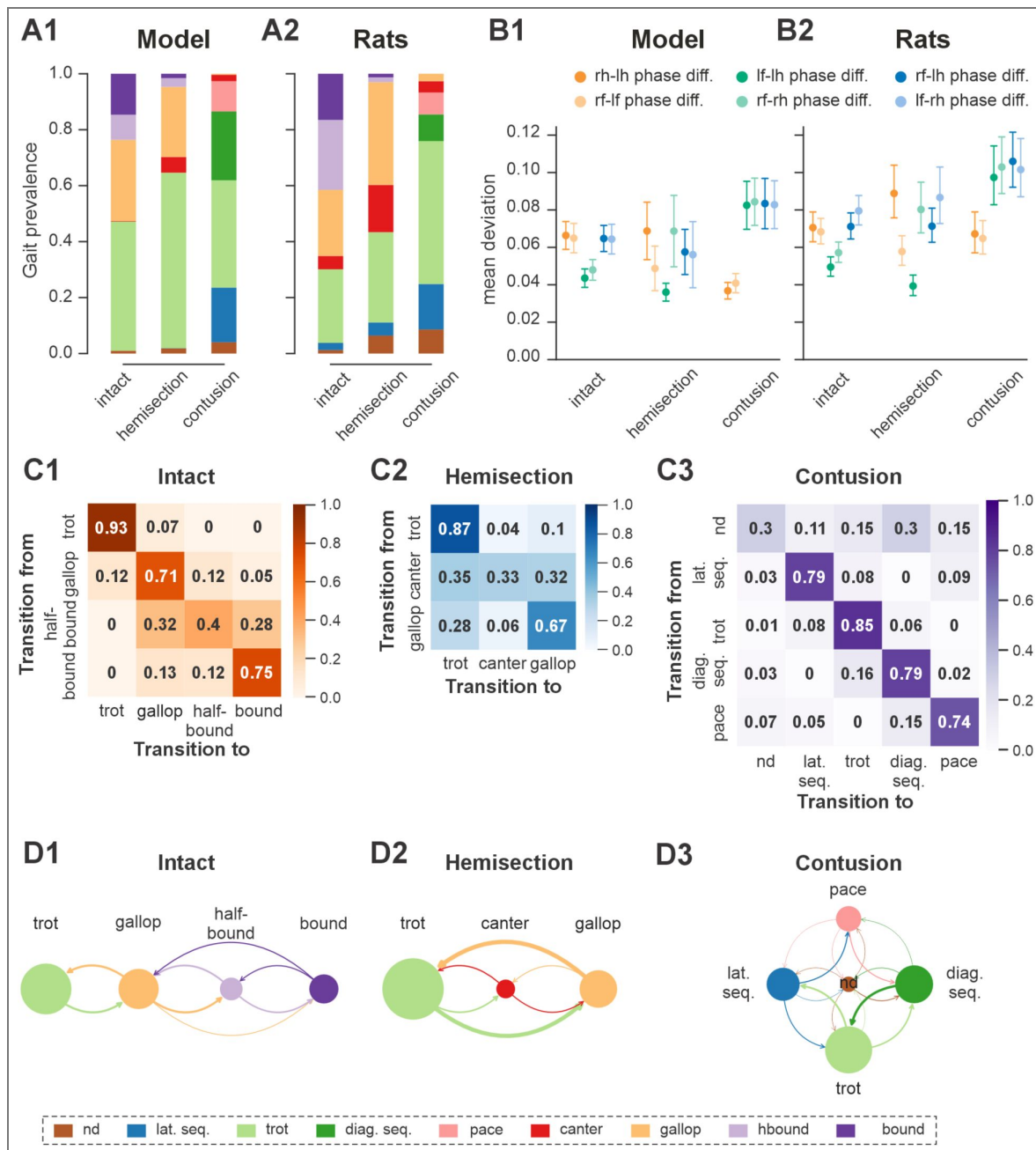


Figure 5. Gait prevalences, variability of interlimb coordination, and gait transition probabilities in model and rats.

A1 Prevalence of gaits in the intact model and following simulated recovery from hemisection and contusion injury. A2 Prevalence of each gait across intact rats and rats after recovery from hemisection and contusion injury (recalculated from Danner et al. (2023) [DOI](#)). B1, B2 Means of the deviations from the moving average of each phase differences for intact case and after recovery from hemisection and contusion injuries for the model (B1) and rats (B2). Detailed statistical results for rats are reported in Danner et al. (2023) [DOI](#). C1–C3 Matrices of gait transition probabilities in the intact model and following recovery from hemisection and contusion injury. D1–D3 Gait transition graphs, where nodes represent gaits (size is proportional to their prevalence) and edges represent gait transitions (line widths are proportional to their frequency of occurrence). l-, left; r-, right; -f, fore RG/limb; -h, hind RG/limb; nd, not-defined; diag., diagonal; lat., lateral; seq., sequence; RG, rhythm generator.

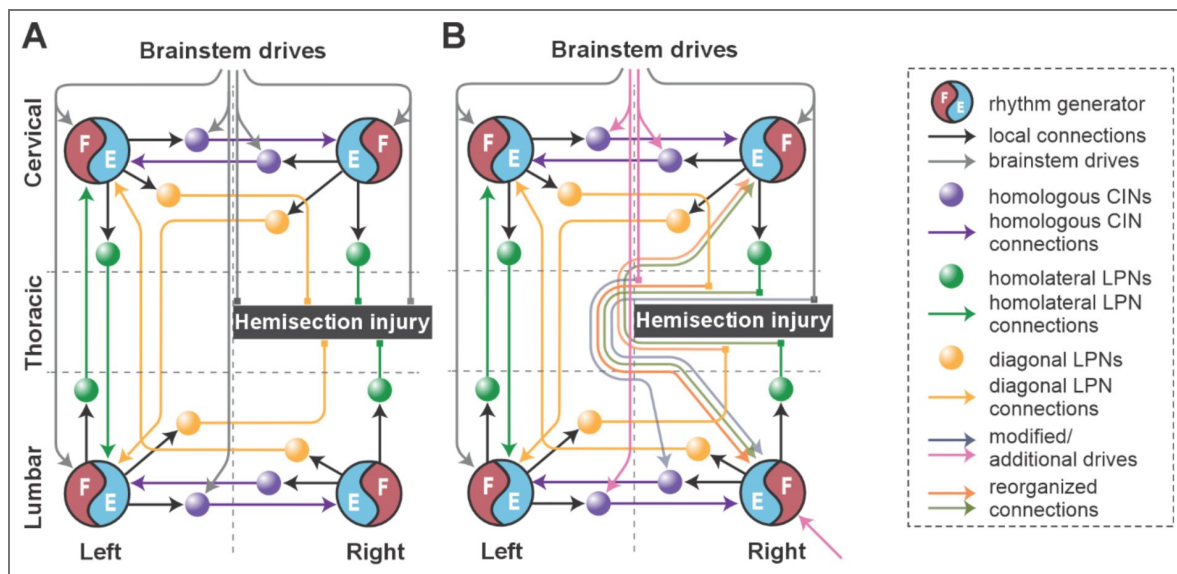


Figure 6. Conceptual schematic of the impact of hemisection injury on the spinal locomotor circuitry (A) and its reorganization after recovery from hemisection predicted by the model (B).

In B, affected long propriospinal neuron (LPN) connections recovered functionally through detour pathways. Drives to lumbar and cervical commissural interneurons (CINs) were altered to strengthen left-right alternation. Drive to the ipsilesional lumbar rhythm generator (pink arrow at the bottom) was substituted by regenerated brainstem input and/or afferent feedback.

- iv. reduction of inhibitory drive to cervical and contralesional lumbar V0_v CINs to the same strength (50%), which through disinhibition strengthened left–right alternation.

The post-hemisection model exhibited frequency-dependent expression of interlimb coordination and gait qualitatively similar to those observed in rats following recovery from hemisection (Figs 5 [5](#), [7](#), and [8](#) and Fig. 2 in Danner et al., 2023 [5](#)). Exemplary bouts of locomotor activities from both the model and animals showed progressive transitions from trot to canter, followed by transverse gallop and back to trot (Fig. 7A1,A2 [5](#), upper panels). In both the model and in rats, hind left–right normalized phase differences during gallop and canter were above 0.5, signifying use of the left-lead variants of both gaits (Fig. 7A1,A2 [5](#), bottom panels). Across all steps, average extensor phase timing and the normalized phase differences for each gait were qualitatively similar between the modeling and experimental results (Fig. 7B1,B2 [5](#)), with the most noticeable difference being the left–right hindlimb phase difference during trot, which in the model showed a stronger asymmetry, i.e., was shifted farther from alternation.

In both the model and in rats, the maximal locomotor frequencies were slightly reduced (compared to the pre-injury case shown in Fig. 4 [5](#)) to approximately 6–6.5 Hz (Fig. 8 [5](#)), and the high-frequency, synchronized gaits (half-bound gallop and bound) were expressed in very few steps (see also Figs 5A1,A2 [5](#) and [8](#)). On the other hand, the occurrence of canter notably increased compared to the intact case (Fig. 5A1,A2 [5](#)) and was distributed across a wide range of generated frequencies (Fig. 8 [5](#)). In both the model and rats, trot (62% of all steps in the model and ~32% in rats) and gallop (~25% in model and ~36% in rats; Fig. 5A1,A2 [5](#)) remained prevalent and were also more stable (high probability of subsequent steps being of the same gait) than canter (Fig. 5C2,D2 [5](#) and Fig. 2H2, I2 in Danner et al. 2023 [5](#)). Canter often occurred as a transitional gait (high probability that next step is of a different gait) and was frequently skipped at transitions from trot to gallop or gallop to trot (Fig. 5C2,D2 [5](#) and Fig. 2H2, I2 in Danner et al. 2023 [5](#)).

The frequency-dependent distribution of phase differences across bouts (Fig. 8A1,B1 [5](#)) exhibits several features consistent with experimental observations in rats following recovery from hemisection injury (Fig. 8A2,B2 [5](#)). All steps were expressed on a continuum of phase differences that appears to be a subset of the phase differences of their pre-injury counterparts (Fig. 4A1,B1 [5](#)). Hind left–right normalized phase differences showed a progressive deviation from alternation (~0.5 phase difference) to quasi-synchronization (phase difference close to 0/1) that occurred only in the direction of the left-lead canter and gallop; right-lead canter and gallop were almost completely lost (Fig. 7C1,C2 [5](#)). Hind left–right synchronization (phase difference close to 0/1; characteristic for half-bound gallop and bound) was almost never reached. Furthermore, there was a left–right asymmetry of the homolateral phase differences: the ipsilesional homolateral normalized phase differences exhibited lower values compared to pre-injury and to the contralesional homolateral phase differences.

Finally, the model exhibited changes in variability of phase differences relative to the pre-injury condition that were similar to those observed in rats (Fig. 5B1,B2 [5](#)). Specifically, compared to pre-injury, the variability (mean deviation from their moving average) of fore left–right and the contralesional (left) homolateral phase differences decreased, whereas variability of the ipsilesional (right) homolateral phase difference increased. The variability of the other phase differences remained at a similar level to their pre-injury values, with a slight increase in the hind left–right phase difference and a slight decrease of the diagonal ones. Although the hind left–right phase difference increased more strongly in rats (Danner et al. 2023 [5](#)), the model results overall are in qualitative agreement with the experiments. It is interesting to note here that the variability of the contralesional (left) homolateral phase difference decreased in the model even though the contralesional LPN connection weights remained at pre-injury levels.

Altogether, these results show that the suggested reorganization of long propriospinal connectivity and descending drive allowed the model to reproduce key features of rat gait expression following recovery from thoracic hemisection injury: expression of a subset of pre-injury gaits, loss of the

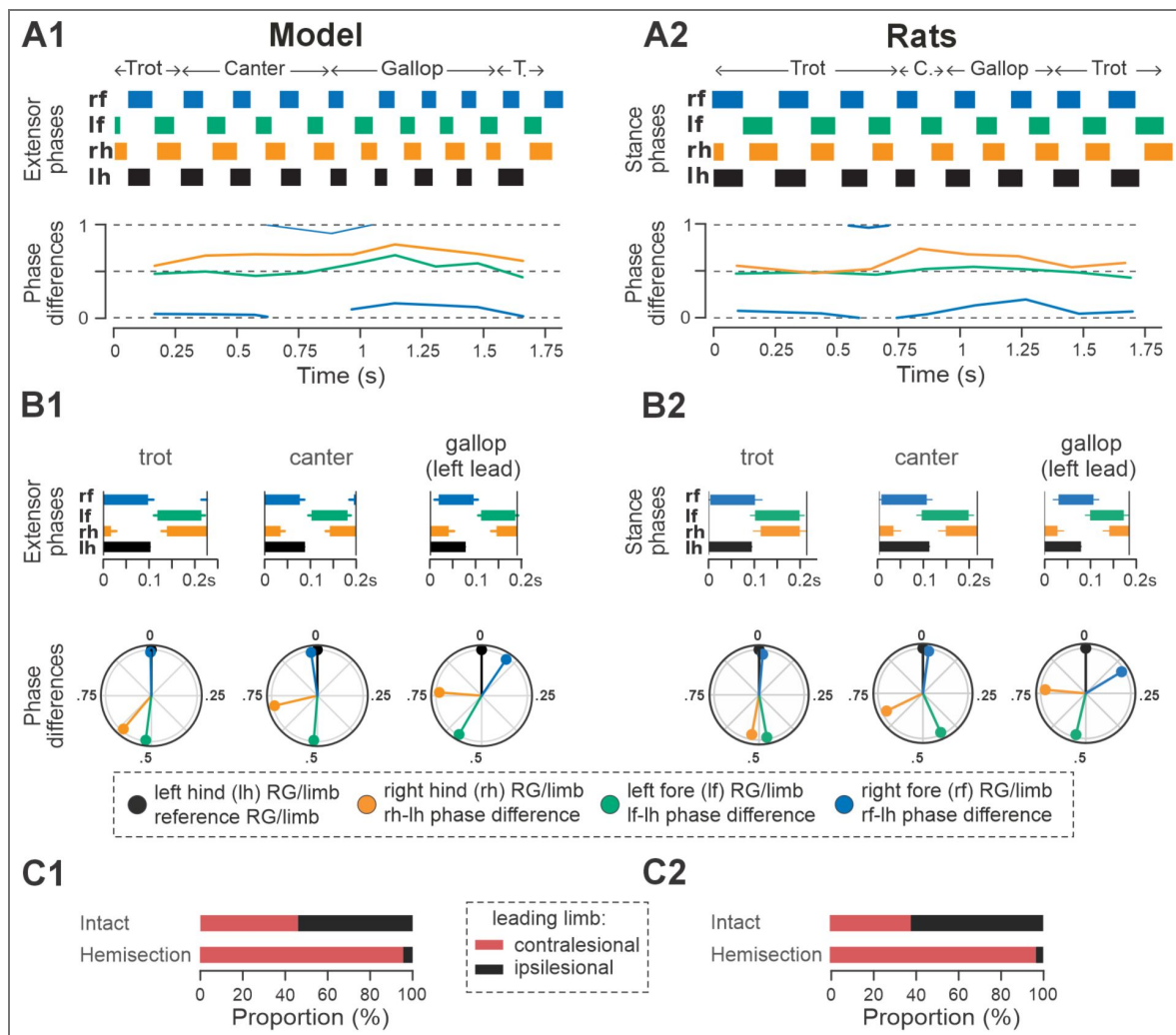


Figure 7. Gait expression in the model and rats following recovery after hemisection.

A1, A2 Extensor/stance phases (upper panels) and instantaneous normalized phase differences (bottom panels) of representative bouts the model (A1) and a rat (A2). B1, B2 Average extensor/stance phases (upper panels) and circular plots of average normalized phase differences for each gait (bottom panels) expressed in the post-hemisection model (B1) and rats (B2). Detailed statistical results for rats are reported in Danner et al. (2023). C1 Prevalences of lead RG in the intact model and following simulated recovery after hemisection for gallop and canter. C2 Prevalence of leading limbs (left or right forelimb that touches down second) pre-injury (intact) and after recovery of hemisection for gallop and canter in rats. Adapted from Danner et al. (2023). T., trot; C., canter; RG, rhythm generator.

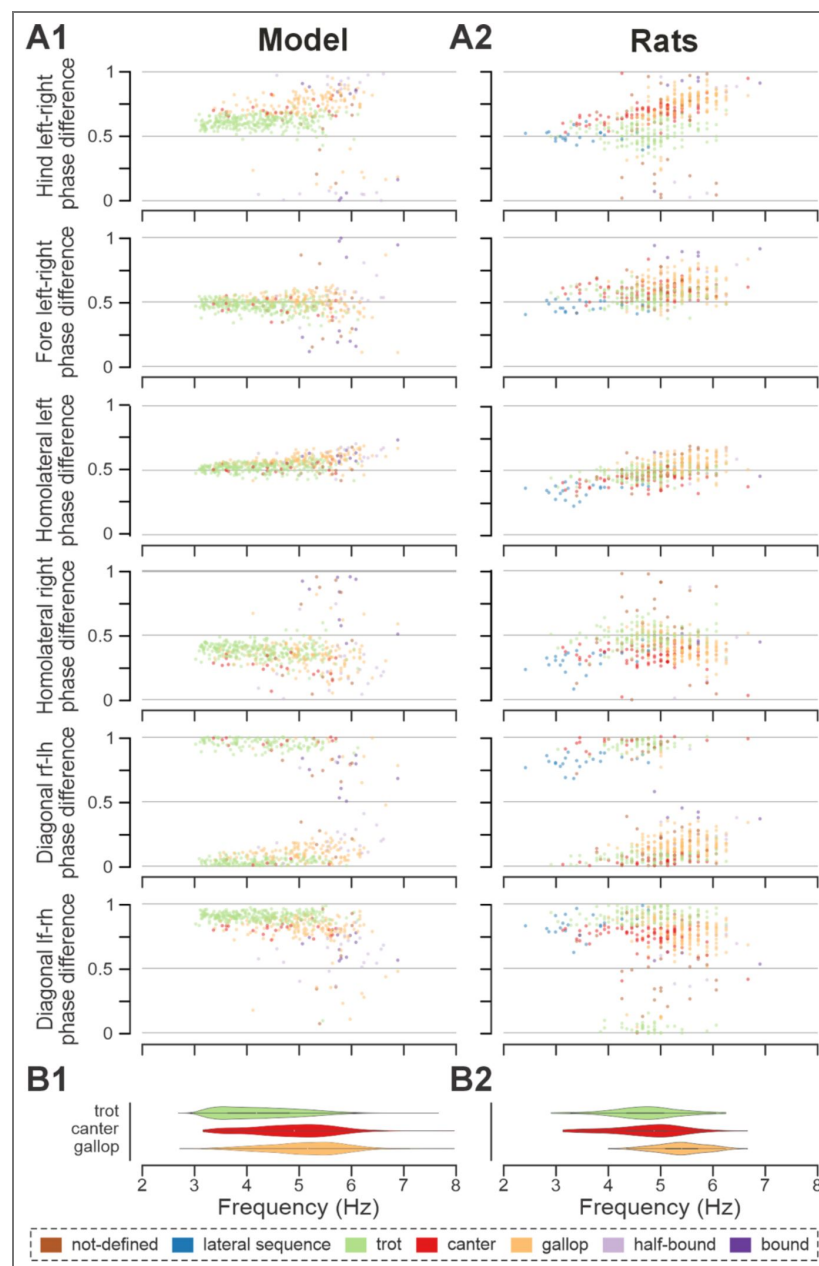


Figure 8. Frequency-dependent distribution of normalized phase differences in the model and in rats following recovery after hemisection.

A1, A2 Scatter plots of normalized phase differences are plotted against frequency of locomotor oscillations. Each dot represents one period/step cycle. Gaits are classified for each period/step cycle and color-coded. B1, B2 Distribution of gaits versus locomotor frequency in the post-hemisection model (B1) and rats (B2). Same number of step cycles are shown for the model and animals; model step cycles were randomly sampled. l-, left; r-, right; -f, fore RG/limb; -h, hind RG/limb; RG, rhythm generator.

highest-speed gaits, and pronounced left–right bias evident in the preferential use of contralesional RG/limb as lead during left–right asymmetric gaits.

Mechanism underlying left–right bias after hemisection

Recovered locomotion following hemisection injury was modeled as a partial (functional) recovery of the pathways severed by the injury and thus, the model included asymmetries of LPN connectivity as well as descending drive (Figs. 1B and 6). To investigate the contributions of these structural asymmetries on the emergence of the left–right bias and loss of high-speed gaits in the post-hemisection model, we created bifurcation diagrams of the normalized phase differences for the brainstem drive parameter α (Fig. 9). These simulations were performed at a low level of noise ($\sigma_{\text{Noise}} = 5$ fA, see Methods) to ensure identification of stable trajectories.

In the intact case, the model generated stable rhythmic locomotor-like activity when parameter α was changed in a stepwise manner from ~ 0.1 to 1.1 (Fig. 9A1). Blue and red lines in Fig. 9A1 indicate the stable phase differences with increase or decrease of α , respectively. Discrepancies between the red and blue lines (when $\sim 0.8 < \alpha < 0.92$) indicate regions of hysteresis and multi-stability. For these α values, multiple stable solutions coexisted and could be expressed depending on the initial conditions. The increased frequency was accompanied by sequential gait changes from lateral-sequence steps to trot to gallop and then half-bound gallop and bound. In the region of multi-stability, both trot and gallop were stable. Importantly, for the left–right asymmetric gaits (gallop and half-bound gallop), where the hind and fore left–right normalized phase differences are not at 0.5 or $0/1$, both left–right symmetric solutions were stable. This corresponds to the stability of gaits with left and right lead RGs. The gait transitions and the corresponding bifurcations in the intact model are similar to our previous mouse models and are described in detail in Danner et al. (2017). The frequency range and step cycle compositions (relative phase durations) for all gaits as well as gait expression were consistent with rat locomotion (Danner et al. 2023).

In the case of simulated hemisection, the bifurcation diagrams show (Fig. 9A2) that the range of brainstem drive parameter α , in which the model is stable, was reduced compared to the intact case. This resulted in a reduction of the maximal frequency to ~ 6 Hz. With increasing drive, the model transitioned from a gait in-between lateral-sequence and canter to trot and then gallop. Half-bound gallop and gallop were lost as stable solutions, and all phase differences lost their second branches. The hind left–right phase difference was shifted above 0.5 (perfect alternation) and progressively increased with increasing α , resulting in continuous transition from trot to gallop only in the direction of left (contralesional) lead sequences. The fore left–right, the contralesional (left) homolateral, and the diagonal phase difference involving the contralesional hind RG all maintained a similar trajectory to their pre-injury counterparts at low α -values before the trot-to-gallop bifurcation. On the other hand, the ipsilesional (right) homolateral and diagonal phase difference involving the ipsilesional hind RG were shifted compared to their pre-injury trajectories.

Next, we investigated how injury-affected LPN connectivity and drive to the ipsilesional hind RG contributed to altered gait expression following hemisection injury. We created bifurcation diagrams for simulations in which only the injury-affected LPN connections (Fig. 9A3) or the drive to the ipsilesional hind RG (Fig. 9A4) were set to their post-injury recovered values (40% and 90% respectively), while all other connection and drive weights were left at their intact values.

Interestingly, the lowered connection weights of the injury-affected and partially recovered LPN weights caused a progressively increasing bias of the left–right phase differences with increasing speed towards ipsilesional (right) lead gallops (Fig. 9A3) — the opposite direction as observed after hemisection. This is followed by a bistable bifurcation (at $\sim 0.85 < \alpha < 0.95$) towards left-lead sequence gallops.

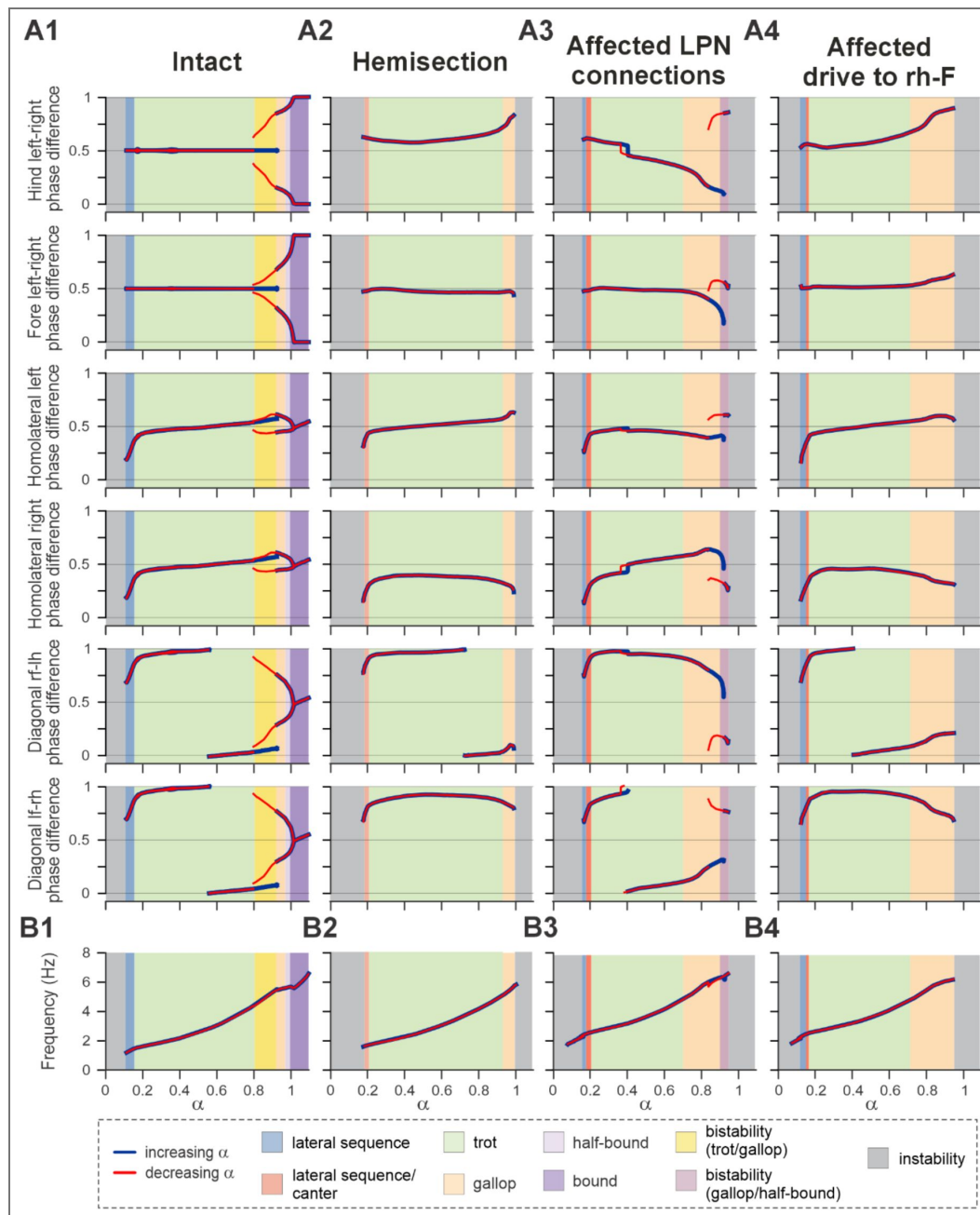


Figure 9.

Bifurcation diagrams of the intact model (A1), following simulated recovery after hemisection (A2), and for model versions where only LPN connections were affected (40% of the pre-injury values; A3) or only brainstem drive to the ipsilesional hind RG was reduced (to 90% of the pre-injury value; A4). Diagrams are plotted against the bifurcation parameter α and with reduced noise, $\sigma_{\text{Noise}} = 5$ fA. Normalized phase differences of 0.5 correspond to alternation, whereas phase differences of 0 or 1 correspond to synchronization. B1–B4 Dependency of frequency of locomotor oscillations on parameter α . Blue and red lines indicate stable phase differences or frequency with stepwise increase and decrease of parameter α , respectively. Colored areas indicate the expressed gait. l-, left; r-, right; -f, fore RG/limb; -h, hind RG/limb; RG, rhythm generator.

On the other hand, the slightly (90% of pre-injury value) lowered drive to the ipsilesional hind RG (Fig. 9A4) caused a left–right asymmetry with a progressive shift away from trot towards gallop and half-bound gallop in the same direction as observed in rats after recovery from hemisection — towards left–lead (contralesional) sequence gaits. Indeed, all six phase differences exhibit trajectories similar to the simulated hemisection (Fig. 9A2), although with stronger deviations of the left–right fore and hind phase differences from alternation at higher frequencies.

Thus, in the model, injury-induced and partially recovered left–right asymmetries of the LPN connections and descending drive to ipsilesional RG have opposing effects on left–right asymmetry of interlimb coordination and gait expression. The hemisection model is a result of the interaction of both of these influences, but the effect of the lower drive to ipsilesional hind RG compared to the other three RGs dominates and largely causes the observed left–right bias.

Modelling recovery from thoracic contusion injury

A midline (left–right) symmetrical thoracic contusion injury (Figs 1C and 10A) partially damages the ascending and descending pathways on both sides of the spinal cord, including supraspinal inputs and long propriospinal connections. However, similar to the hemisection injury, the cervical and lumbar circuitry including their homologue pathways remain intact. After a moderate thoracic midline contusion injury, rats regain functional locomotion dominated by left–right alternating gaits (Danner et al. 2023), and even slower maximum speed than after hemisection. They lose the ability to perform non-alternating gaits (gallops and bounds) and adopt atypical gaits not observed in uninjured animals, such as diagonal-sequence and pace. These two novel gaits are characterized by left–right alternation of both the forelimbs and hindlimbs. The diagonal sequence is a four-beat gait pattern where a hindlimb step is followed by a step of the contralateral (diagonal) forelimb, rather than the ipsilateral forelimb, as is the case in the lateral sequence. Pace is a two-beat gait in which the homolateral limbs move in synchrony. Previously, we hypothesized that the post-contusion gait changes are a result of weakened fore–hind coupling together with maintenance of left–right alternation (Danner et al. 2023). We assumed that after symmetrical thoracic contusion injury the brainstem drive and ascending and descending LPN connections on both sides of the model are significantly damaged (Figs 1C and 10A). Thus, to model circuit reorganization underlying recovered post-contusion locomotion, we assumed that spared sublesional lumbar circuitry undergoes plastic reorganization and that supraspinal control over the intact supraspinal cervical circuitry is altered. Then, by manually tuning model parameters, we looked for possible changes of connection weights in the model that result in reproduction of locomotor function of rats following recovery from contusion, while also keeping the connection weights of the injured pathways low. Note that we did not make any assumption about the proportion of the translesional fibers that remain functionally intact.

This process resulted in the following specific suggestions that we implemented in the model:

- i. reduced brainstem drive to the sublesional lumbar RGs was substituted with enhanced afferent input and/or changes in their intrinsic excitability (simulated as additional drives to the hind RGs);
- ii. descending control over the supraspinal cervical circuitry via (tonic) brainstem drive was adjusted so that the oscillation frequency of the cervical RGs matched those of the lumbar RGs (simulated by equal drives to cervical and lumbar RGs) and to secure fore left–right alternation (reduction of inhibitory drives to cervical V0 CINs);
- iii. commissural pathways in the lumbar compartment were reorganized to secure hind left–right alternation (descending drive to V0_l CINs was removed, and their excitability was altered by an additional constant drive) (Fig. 10B); and
- iv. finally, connection weights of all inter-enlargement LPN connections were set to 5% of their pre-injury values.

An exemplary bout produced by the post-contusion model is shown in Fig. 11A1. The model progressively transitioned from a diagonal sequence gait to trot then across lateral sequence to pace and finally back to the diagonal-sequence gait. The example bout of rat post-contusion recovery (Fig. 11A2) exhibited the same sequence of gait transitions except that it starts

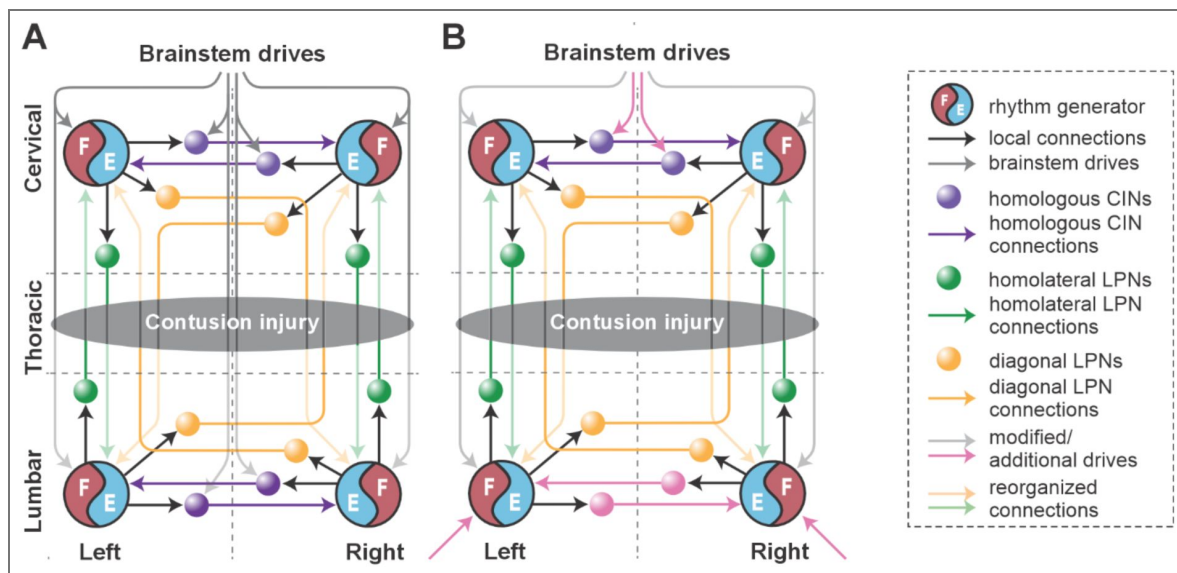


Figure 10. Conceptual schematic of the contusion injury (A) and following recovery after contusion in the model (B).

The weights of the long propriospinal neuron (LPN) connections between the cervical and lumbar compartments were significantly reduced. Brainstem drive to the lumbar rhythm generators (RGs) was substituted with additional drives to these RGs (pink arrows at the bottom). Brainstem drives to the cervical RG were adjusted to match oscillation frequency of lumbar RGs (gray arrows at the top). Inhibitory drives to cervical V0v commissural interneurons (CINs) were reduced to secure fore left-right alternation (pink arrows at the top). Commissural pathways in the lumbar compartment were reorganized to secure hind left-right alternation (pink arrows at the bottom).

and ends with a lateral sequence gait pattern. Note that fore and hind left–right alternation was maintained across all steps of the bouts, while the homolateral and diagonal phase differences progressively drift in parallel across the full cycle reaching their approximate initial values at the end of the bouts (Fig. 11A1,A2 [↗](#), bottom panels). In both cases, the fore limbs/RGs performed one more step cycle than the hind limbs/RGs (Fig. 11A1,A2 [↗](#), upper panels).

Across simulated bouts, the post-contusion model expressed lateral and diagonal-sequence gait patterns, as well as trot and pace (Fig. 5A1 [↗](#)). Like contused rats (Fig. 5A2 [↗](#); Danner et al. 2023 [↗](#)), the model did not express non-alternating gaits (gallop, half-bound gallop and bound) but produced left–right alternating gaits that the intact model did not (including pace and the diagonal-sequence pattern; Fig. 11B1 [↗](#)). The maximal locomotor frequencies were reduced up to approximately ~6 Hz (Fig. 12 [↗](#)) and average gaits of the model exhibited similar extension phase patterns and phase differences to those observed in rats (Fig. 11B1,B2 [↗](#)). Across the expressed range of frequencies, both fore and hind left–right normalized phase differences remained alternating (around 0.5) — which is in contrast to the intact case where fore and hind left–right phase differences progressively transition from alternating to in-phase synchronization with increasing frequency (Fig. 12A1,A2 [↗](#)). On the other hand, all fore–hind phase differences (both homolateral and diagonal phase differences) were highly variable and distributed across the whole cycle. Frequency-dependence of gait expression was largely lost, with all gaits expressed across the full range of frequencies (Fig. 12A1,A2 [↗](#)). Yet, in the model diagonal-sequence steps were more frequent at low frequencies and lateral-sequence steps at high frequencies. This is different to contused rats, which expressed these gaits on average at comparable frequencies (Fig. 12B1,B2 [↗](#)).

Interestingly, gait transition probabilities in the post-contusion model showed a clear pattern (Fig. 5C3,D3 [↗](#)) and apart from the not-defined category (capturing all other gaits), the expressed gaits were relatively stable: a step of one gait was likely to be followed by another of the same gait (all probabilities 0.74–0.85). Transitions predominantly occurred between neighboring gaits along the sequence: lateral-sequence, trot, diagonal-sequence, pace, and back to lateral-sequence (Fig. 5C3,D3 [↗](#)). The idealized versions of these gaits are evenly distributed along a line (or rather circle) in phase space (defined by the three orthogonal phase-differences: hind left–right, homolateral, and diagonal). The line connects trot and pace in both directions (through lateral-sequence in one and diagonal-sequence in the other direction) and represents the only remaining degree of freedom, assuming that fore and hind left–right alternation was maintained. This organization also explains the experimentally observed distribution of phase differences in contused rats (Danner et al. 2023 [↗](#)).

These results are further complemented by the analysis of phase difference variability (Fig. 5B1,B2 [↗](#)). Specifically, the mean variability (mean deviation from the moving average) of both fore and hind left–right phase differences were lower compared to the variability of all fore–hind phase differences in both the model and in rats. Comparing variability post-contusion to the pre-injury condition revealed some discrepancies between the model and experimental results. In the model, variability of fore and hind left–right phase differences decreased after contusion, whereas the experimental results showed no significant change. This suggests that the model may have overestimated the strengthening of left–right coupling due to disinhibition of cervical V0_v CINs and reorganization of the lumbar circuitry. However, the mean variability of all fore–hind phase differences (left and right homolateral and diagonal) increased in both the model and experimental results.

Overall, the model supports our previous hypothesis that, following a symmetrical moderate thoracic contusion injury coupling between the left and right RGs within each enlargement remained at (or recovered back to) pre-injury levels, while the coupling between the lumbar and cervical circuits was significantly reduced. These results suggest that plasticity with the local circuitry linking the left and right RGs within each enlargement are critical for recovery following a symmetrical thoracic contusion injury.

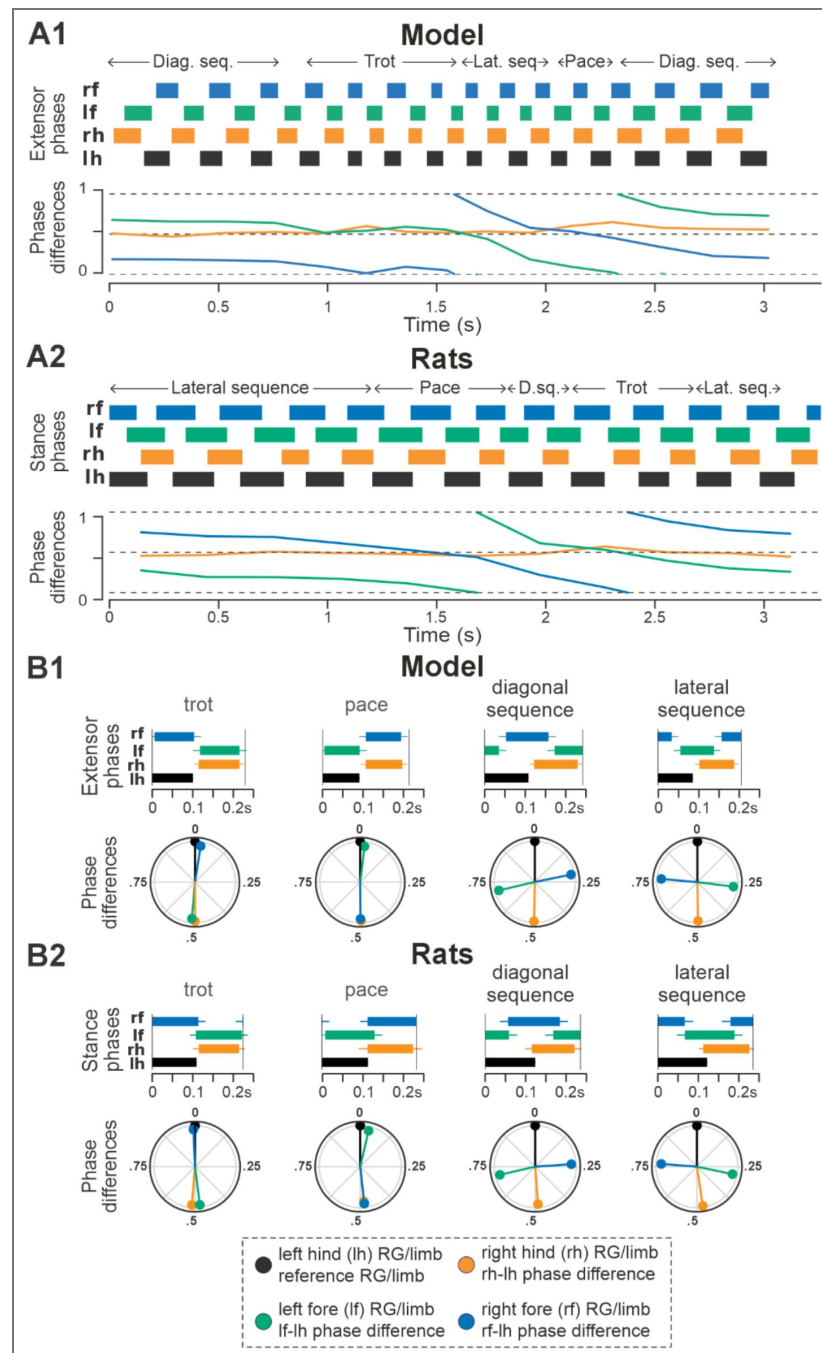


Figure 11. Gait expression in the model and rats following recovery after contusion.

A1, A2 Extensor/stance phases (upper panels) and instantaneous normalized phase differences (bottom panels) of representative bouts the model (A1) and a rat (A2). B1, B2 Average extensor/stance phases for each gait (upper panels) and circular plots of average normalized phase differences for each gait (bottom panels) expressed in the post-contusion model (B1) and rats (B2). Detailed statistical results for rats are reported in Danner et al. (2023). D.sq., diag. seq., diagonal-sequence; lat. seq., lateral-sequence; RG, rhythm generator.

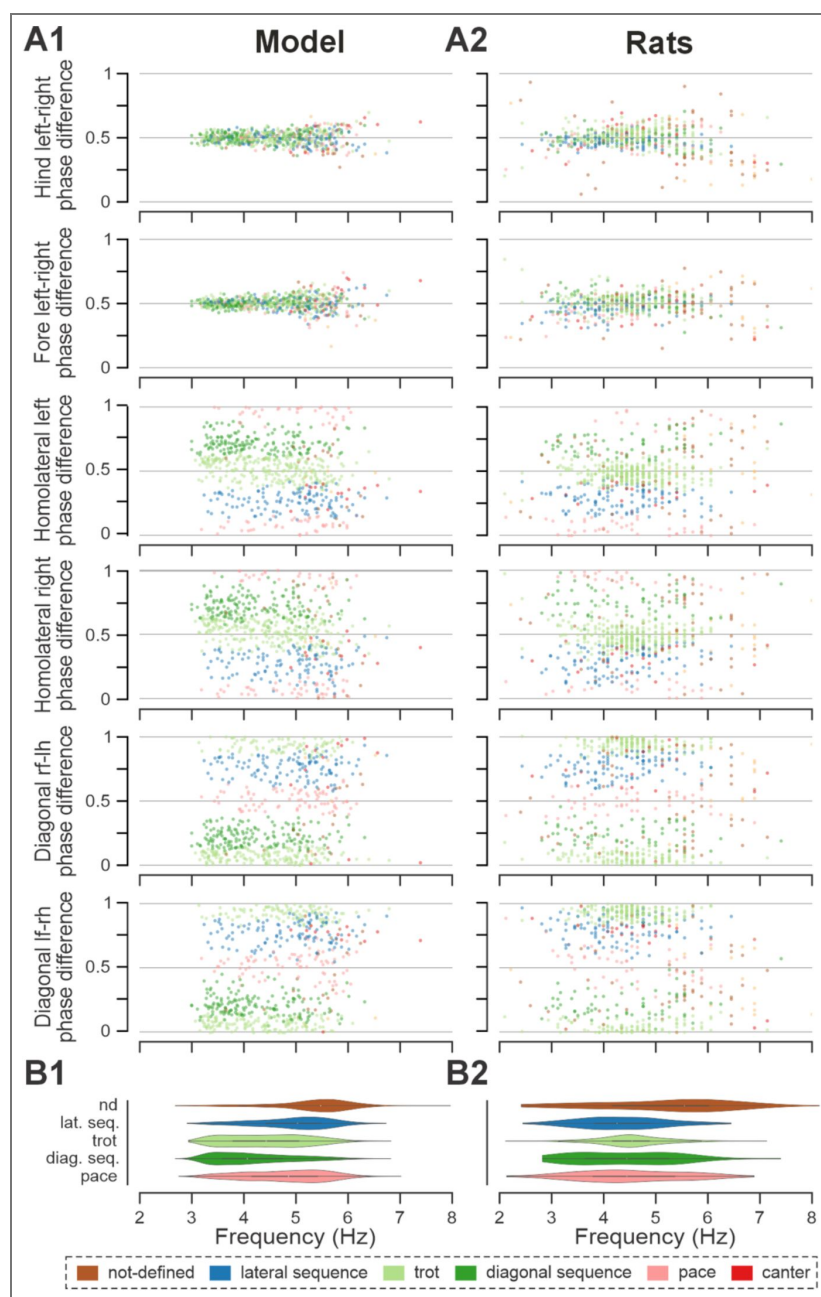


Figure 12. Frequency-dependent distribution of normalized phase differences in the model (A1) and in rats (A2) following recovery after contusion.

A1, A2 Scatter plots of normalized phase differences are plotted against frequency of locomotor oscillations. Each dot represents one period/step cycle. Gaits are classified for each period/step cycle and color-coded. B1, B2 Distribution of gaits vs. locomotor frequency in the model (B1) and rats (B2). l-, left; r-, right; -f, fore RG/limb; -h, hind RG/limb; RG, rhythm generator.

Sensitivity of hemisection and contusion models to connectivity changes

To evaluate which connectivity changes most strongly shaped speed-dependent locomotor behavior after injury, we performed a local sensitivity analysis of the contusion and hemisection models. This analysis quantified how small variations (in the range of [80%,125%]) in the connection weights of key pathways affect post-injury frequency-dependent gait expression. The outcome measure was the Earth Mover's Distance (EMD; Rubner et al., 1998) of each perturbed simulation from the corresponding post-injury baseline model in a 7-dimensional feature space (six interlimb phase differences plus locomotor frequency). Sensitivity was expressed as Sobol indices (Fig. 13), which quantify the fraction of variance in the EMD attributed to each parameter (Sobol', 2001). High indices identify parameters whose perturbations most strongly alter frequency-dependent gait expression (for more details see Methods).

In the hemisection model, drive to the ipsilesional hind/lumbar flexor rhythm generator center was the dominant determinant of locomotor outcomes (Fig. 13A), with both first- and total-order indices reaching ~0.8–0.9. Varying this drive within the sensitivity range produced large, systematic shifts in phase-difference–frequency relationships and gait distributions (Fig. 13—figure supplement 1A,B), consistent with its high Sobol values. Lumbar commissural interneuron (V0 and V3) connection weights were the only other parameters with appreciable influence (total-order indices ~0.1), and their variation also produced clear changes in phase differences and gait patterns (Fig. 13—figure supplement 1D,E). In contrast, long propriospinal pathways and cervical CINs showed very low sensitivity indices (≤ 0.01); for example, recovered LPN connection weights resulted in outputs nearly indistinguishable from the baseline model (Fig. 13—figure supplement 1A,C), representative of other low-sensitivity parameters.

A similar pattern was observed in the contusion model. Drive to the sublesional lumbar flexor rhythm generator centers dominated the sensitivity profile (total-order 0.96; first-order 0.69; Fig. 13B), and varying this drive generated robust changes in frequency-dependent distributions of phase differences and gaits (Fig. 13—figure supplement 2A,B). Lumbar V0 and V3 CIN connection weights were the next largest contributors (total-order 0.08–0.22; Fig. 13B), and their variation produced clear, albeit smaller, distortions in phase relationships and gait expression (Fig. 13—figure supplement 2D,E). In contrast, cervical CINs and ascending/descending LPN pathways had negligible effects on locomotor output (total-order indices < 0.02), with example simulations of varying ascending LPN weights closely resembling baseline behavior (Fig. 13—figure supplement 2A,C).

Partial dependence plots further revealed how interactions between lumbar V0 and V3 commissural pathways shape locomotor activity in both hemisection and contusion models (Fig. 13—figure supplement 3). The large difference between total-order and first-order sensitivity for these pathways (Fig. 13A,B) indicates that their influence arises mainly through interactions rather than isolated changes to either V0 or V3 pathways. Consistent with this, covarying V0 and V3 together (either increasing or decreasing both within the sensitivity range) kept predicted EMD low, indicating that coordinated scaling preserves overall locomotor behavior. In contrast, disrupting the balance between the two pathways (by changing only one, or by altering them in opposite directions) produced substantially larger deviations from the baseline pattern. These results suggest that the relative balance of V0 and V3 connectivity, rather than their absolute strengths individually, is critical for maintaining post-injury gait expression in the model.

Taken together, the results demonstrate that across both injury models, two classes of parameters substantially influence frequency-dependent gait expression: (1) drive to the sublesional lumbar rhythm generators and (2) lumbar commissural (V0/V3) interneuron connectivity, whose impact depends critically on their coordinated balance. Within the tested perturbation range, changes in LPN pathways and cervical CINs had only minor effects on model output. This emphasizes the central role of restoration of activation of sublesional lumbar rhythm-generating circuits and strengthened commissural pathways in shaping post-injury locomotor patterns.

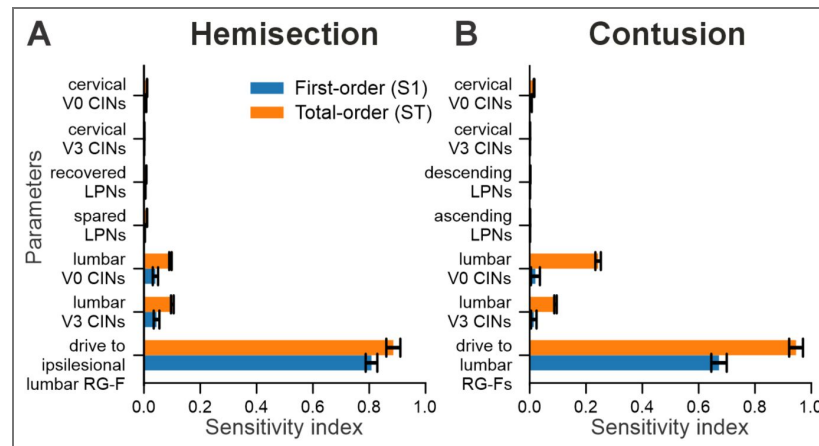


Figure 13. Local sensitivity analysis of hemisection (A) and contusion (B) models.

Shown are Sobol first-order (S1) and total-order (ST) sensitivity indices quantifying how small parameter variations (80–125% of baseline) affected locomotor behavior, measured as the Earth Mover’s Distance (EMD) across six interlimb phase differences and locomotor frequency. Parameters include drive to the sublesional lumbar flexor rhythm generator centers (RG-F; ipsilesional in hemisection, bilateral in contusion), connection weights of cervical and lumbar V0 and V3 commissural interneuron (CIN), and long propriospinal neuron (LPN) pathways. Error bars indicate 95% confidence intervals.

Discussion

The brain and spinal cord have a high capacity to adapt and reorganize. Indeed, following incomplete spinal cord injuries, neural plasticity occurs across the breadth of the nervous system (Filli and Schwab, 2015). This includes reorganization of supraspinal motor centers, compensatory structural and functional plasticity of spared descending, propriospinal, and sensory neurons above and below the lesion (Ballermann and Fouad, 2006; Rosenzweig et al., 2010; Filli et al., 2014; Takeoka et al., 2014; Zörner et al., 2014), local regrowth/sprouting of transected fibers, as well as intrinsic changes to sublesional spinal interneurons and motoneurons (Barriere et al., 2008; Cohen-Adad et al., 2014; Takeoka and Arber, 2019). In this computational modeling study, we have investigated potential reorganization strategies of neural connectivity underlying recovery of locomotor function following thoracic lateral hemisection and moderate midline contusion injuries in rats, representing asymmetrical and symmetrical injuries, respectively, that exhibit roughly comparable functional recovery (Danner et al., 2023). The model itself represents neuronal populations resident in the cervical and lumbar enlargements with axons of ascending and descending inter-enlargement propriospinal neurons along with reticulospinal drive to the individual hind and fore RGs. The model reproduced key features of speed-dependent gait expression of rats before and after spinal cord injury. It suggests that recovery after lateral hemisection involves partial functional restoration of descending drive and long propriospinal pathways, whereas recovery following midline contusion relies on reorganization of sublesional lumbar circuitry combined with altered descending control of cervical networks. Across both injuries, sensitivity analyses consistently identified two principal determinants of recovered locomotion: restored excitability of the sublesional lumbar rhythm-generating circuits and appropriate connectivity and balance of lumbar V0/V3 commissural interneuron pathways. These findings highlight the central role of sublesional lumbar rhythm-generating and commissural circuitry in shaping post-injury locomotor function.

Descending drive from the reticular formation controls locomotor speed and gait in intact animals (Capelli et al., 2017; Ausborn et al., 2019; Hsu et al., 2023) and substantial structural and functional plasticity of reticulospinal neurons occurs in parallel with recovery of locomotor function following hemisection injury (Ballermann and Fouad, 2006; Filli et al., 2014; Engmann et al., 2020; Lemieux et al., 2024). In our model, drive from the reticular formation controls the oscillation frequency of each rhythm generator and modulates the activity of commissural interneurons controlling interlimb coordination. The model suggests that following the asymmetric lateral hemisection injury, the drive to the ipsilesional hind rhythm generator recovers to around 90% of its pre-injury strength (Fig. 6B). The strength of this drive reflects the net excitatory input to the population of flexor rhythm generator neurons and cannot be easily interpreted in terms of the number of regenerated axons or synaptic terminals. Thus, while these results suggest substantial reorganization of reticulospinal interactions with the lumbar circuitry, other factors, such as changes in intrinsic neuronal properties, increased prevalence of excitatory connections between constituent neurons, and strengthened afferent input (Barriere et al., 2008; Cohen-Adad et al., 2014; Takeoka et al., 2014; Takeoka and Arber, 2019; indicated by the pink arrow to the ipsilesional hind RG in Fig. 6B) can also affect the excitability of the sublesional rhythm generator neurons and help compensate for the reduced descending drive.

Inter-enlargement long propriospinal neurons couple the cervical circuits controlling the forelimbs with the lumbar circuits controlling the hindlimbs and are essential for interlimb coordination (Figs 1A and 2; Edinger, 1896; Nathan and Smith, 1959; Dutton et al., 2006; Ruder et al., 2016; Flynn et al., 2017; Frigon, 2017). These neuronal populations are roughly 50% ipsilaterally and 50% contralaterally projecting with the axons of the latter crossing the midline at the level of the cell bodies (Reed et al., 2006; Pocratsky et al., 2020; Shepard et al., 2023). Lateral thoracic hemisection injury severs all these axons unilaterally (Fig. 6A). Similar to the predicted recovery in descending drive, the model also predicts that partial (functional) recovery of the severed long propriospinal pathways is helpful and possibly necessary for recovery (Fig. 6B). This recovery was simulated by setting the connection weights of the injured long propriospinal neuron populations and drive connections to a proportion of their pre-

injury strengths. Long-distance regeneration of central axons is unlikely to occur in adult animals (Filli and Schwab, 2015), thus reorganization of these pathways likely depends on local axon growth and synaptic rewiring onto propriospinal neurons with intact axons, resulting in formation of detour pathways through the intact hemicord (Courtine et al., 2008), as well as compensatory mechanisms of spared reticulospinal and long propriospinal neurons, including increased innervation of the ipsilesional lumbar hemicord via sprouting (Ballermann and Fouad, 2006; Courtine et al., 2008; Takeoka et al., 2014; Filli and Schwab, 2015). Indeed, recovery of locomotor function after hemisection injury has been shown to result in an increased prevalence of long propriospinal neurons bypassing the injury by crossing the midline twice and of reticulospinal connections to the ipsilesional lumbar cord (Takeoka et al., 2014). The work by Reed (Reed et al., 2006) uncovered evidence for a small number of descending inter-enlargement neurons with axons that crossed the midline close to the cell body and then again in the lumbar enlargement to terminate ipsilaterally in uninjured animals, suggesting that a descending detour pathway may already exist, even if small. Furthermore, potentiation of ipsilesional muscle responses to optogenetic stimulation of the reticular formation has also been linked to locomotor recovery (Lemieux et al., 2024).

Recovery from lateral thoracic hemisection results in distinct left–right asymmetries of locomotor behavior (Danner et al., 2023). These asymmetries are most obvious at higher speeds and for non-alternating gaits — animals almost exclusively use the contralesional limb as the lead limb. We investigated the individual contribution of the asymmetries of long propriospinal connections and of the descending drive of the simulated hemisection (Fig. 9A3,A4). Interestingly, in the case where only the asymmetrical drive to the lumbar rhythm generators (90% of pre-injury strength to ipsilesional RG) was included and all other connections were set to their pre-injury values, the model (Fig. 9A4) very closely reproduced frequency-dependent changes of interlimb coordination and gait expression of hemisected animals. On the other hand, the asymmetry introduced by the partially recovered long propriospinal connectivity resulted in gait asymmetries in the opposite direction (ipsilesional limb used as lead limb for non-alternating gaits; Fig. 9A3). Thus, the asymmetry of the drive to the ipsilesional hindlimb rhythm generator compensated for the asymmetry of the long propriospinal connections and was the main contributor to the observed gait asymmetries (Figs 7, 8 and 9A2).

In the case of the midline, symmetrical contusion injury, our model suggests that connection weights of 5% of their pre-injury values for the long propriospinal neurons are sufficient to reproduce the experimentally observed changes in interlimb coordination and gait expression. These values are much lower than anticipated based on the proportion of axons (white matter) spared by this type of contusion injury, which affects approximately 70–80% of the overall white matter (Basso et al., 1996) but presumably a much smaller proportion of the axons in the outermost layers of the ventrolateral funiculus where the long propriospinal tracts are located (Basso et al., 1996; Reed et al., 2006; Brown et al., 2024). Note that we assumed that post-contusion recovery, descending drive to cervical rhythm generators is tightly controlled to match the level of excitability of the lumbar rhythm generators. This assumption was modeled by adjusting the drive so that the oscillation frequency of cervical and lumbar rhythm generators match. In reality, the excitability of the lumbar rhythm generators would be affected by somatosensory afferent input and body biomechanics (Rossignol et al., 2006; Nishikawa et al., 2007; Akay et al., 2014; Takeoka et al., 2014; Takeoka and Arber, 2019; Ausborn et al., 2021; Kim et al., 2022). We introduced noise to all neurons to simulate step-to-step variability, but we did not consider afferent feedback or limb/body biomechanics in the model. Thus, we cannot exclude that the long propriospinal connection strengths necessary to reproduce experimental interlimb coordination were underestimated.

Yet, there is experimental support for plasticity resulting in decreased long propriospinal connectivity. Specifically, using viral-based tracing, we recently found that the number of axons/terminals in the cervical enlargement arising from ascending long propriospinal neuron was significantly reduced after recovery from a mild contusion injury (Brown et al., 2024), and further that silencing of a proportion of either ascending or descending long propriospinal

neurons after moderate contusion injury actually improves (normalizes) interlimb coordination and other key locomotor indices (Shepard et al., 2021 [DOI](#), 2023 [DOI](#)). Thus, long propriospinal neurons seem to have a disruptive effect on interlimb coordination after injury. One explanation might be that the injury affects different classes of long propriospinal neurons (ascending vs. descending, homolateral vs. diagonal, left vs. right) differently, leading some to make new connections that are maladaptive to recovery and that (further) reduction of their connectivity via silencing reduces their influence and thereby improves interlimb coordination. In support of this idea, in Brown et al. (2024) [DOI](#), we also found that ascending propriospinal innervation of segments just caudal to the thoracic injury was increased in the same animals where input to C6 was decreased, all relative to uninjured animals. Thus, unguided plasticity of long propriospinal neurons post-injury seems to have a disruptive effect on interlimb coordination.

Silencing of ascending or descending long propriospinal neurons in intact mice or rats has the opposite effect compared to following contusion; it disrupts interlimb coordination in a speed and context dependent manner, causing transient periods of stepping with disordered left–right coordination (Ruder et al., 2016 [DOI](#); Pocratsky et al., 2020 [DOI](#)). Our previous models attributed this effect to the excitatory diagonally projecting long propriospinal neurons (ascending V3 LPNs, descending V0_v LPNs), which ensure diagonal synchronization for trot but also promote left–right alternation because of the tightly controlled homolateral fore–hind alternation by ipsilaterally projecting long propriospinal neurons (Danner et al., 2017 [DOI](#); Ausborn et al., 2021 [DOI](#); Zhang et al., 2022 [DOI](#)). Thus, the influence of long propriospinal interneurons promoting left–right alternation of cervical and lumbar rhythm generators is reduced following contusion injury. Indeed, somewhat more severe contusion injuries that remove more than 80% of white matter result in an increased prevalence of disrupted left–right alternation, and other persistent problems such as dorsal stepping and the loss of the 1:1 forelimb/hindlimb relationship during stepping (Shepard et al., 2021 [DOI](#), 2023 [DOI](#)). Yet, recovery from moderate contusion injuries — where plantar stepping and a 1:1 forelimb-to-hindlimb ratio are recovered, as modeled here (Danner et al., 2023 [DOI](#)) — results in well-controlled left–right alternation of both fore and hindlimbs. Even animals that show the poorest recovery regain stepping, and the number of steps with appropriate left–right alternation significantly exceed the number of disrupted steps (Shepard et al., 2021 [DOI](#), 2023 [DOI](#)). Together, these findings suggest that reorganization and/or altered modulation of intra-enlargement commissural interneurons occurs during recovery to strengthen left–right alternating influences. In the model this was implemented by reduced descending inhibition of cervical V0 commissural interneurons and altered excitability of lumbar V0 commissural interneurons. This specific implementation reflects one possible scenario, as assumed by the model, where V0 commissural interneuron excitability is regulated by descending inhibition. Yet, descending excitation (Danner et al., 2016 [DOI](#)) or descending neuromodulatory influences — such as serotonergic, noradrenergic, or dopaminergic input — may also contribute (Butt and Kiehn, 2003 [DOI](#); Jordan et al., 2008 [DOI](#); Abbinanti and Harris-Warrick, 2012 [DOI](#); Abbinanti et al., 2012 [DOI](#); Perrier and Cotel, 2015 [DOI](#)). In addition, local excitatory V2a (Crone et al., 2009 [DOI](#)) and local inhibitory interneurons contribute to left–right coordination (Kjaerulff and Kiehn, 1997 [DOI](#); Butt and Kiehn, 2003 [DOI](#); Shevtsova et al., 2015 [DOI](#); Danner et al., 2016 [DOI](#), 2017 [DOI](#); Kiehn, 2016 [DOI](#)) and may be modulated by descending input, making them potential targets for reorganization following injury.

Model limitations and future direction

Our model has several limitations. Most importantly, it does not incorporate afferent feedback or body and limb biomechanics, both of which play a major role in both pre- and post-injury locomotion and can affect interlimb coordination (Forssberg et al., 1980 [DOI](#); Rossignol et al., 2006 [DOI](#); Nishikawa et al., 2007 [DOI](#); Aoi et al., 2013 [DOI](#); Akay et al., 2014 [DOI](#); Takeoka et al., 2014 [DOI](#); Takeoka and Arber, 2019 [DOI](#); Kim et al., 2022 [DOI](#); Molkov et al., 2024 [DOI](#)). Some of the connections in our model might indeed be mediated by afferent feedback: for example, ascending long propriospinal neurons receive afferent feedback (Lloyd and McIntyre, 1948 [DOI](#); Flynn et al., 2017 [DOI](#); Frigon, 2017 [DOI](#); Laflamme et al., 2023 [DOI](#)), and commissural interneurons integrate afferent input from various modalities (Edgley et al., 2003 [DOI](#); Jankowska, 2008 [DOI](#); Jankowska et al., 2009 [DOI](#);

Laflamme et al., 2023 [↗](#)). Furthermore, we did not explicitly model reorganization of afferent feedback connectivity (Takeoka et al., 2014 [↗](#); Takeoka and Arber, 2019 [↗](#)). In reality, restoration of sublesional rhythm generator activation may arise not only from partial recovery of descending pathways but also from enhanced somatosensory feedback that partially compensates for lost drive (Forssberg et al., 1980 [↗](#); Lovely et al., 1986 [↗](#); Barbeau and Rossignol, 1987 [↗](#); Takeoka et al., 2014 [↗](#); Frigon, 2017 [↗](#); Takeoka and Arber, 2019 [↗](#)); in the present model, these contributions are lumped together and represented only as changes in effective drive to the rhythm generators. Moreover, we note that the reorganizations presented here represent solutions that are sufficient to reproduce experimentally observed post-injury gait expression and alternative circuit configurations may likewise support recovery. Finally, because our modeling framework identified circuit reorganizations that successfully reproduced the observed post-injury speed-dependent gait expression, it primarily reveals reorganizations that support functional recovery and does not capture potentially maladaptive changes that impair locomotor performance.

Future work will require neuromechanical models that explicitly incorporate phasic afferent feedback, its reorganization after injury, and its interaction with body and limb biomechanics. Experimentally, a systematic characterization of post-injury reorganization in lumbar circuits and their interacting pathways (including supraspinal, long propriospinal, commissural, and afferent inputs, with particular emphasis on those that restore activation of sublesional rhythm generators) is needed, as these elements emerged as central to recovered function in our model. Constraining and validating extended models with such data will be critical for distinguishing circuit reorganizations that promote locomotor recovery from those that are maladaptive and limit it.

Conclusion

In conclusion, challenging animals to perform overground locomotion across the full range of speeds before and after two distinct spinal cord injury models revealed injury-specific deficits in interlimb coordination and gait expression (Danner et al., 2023 [↗](#)). These deficits differed between the two injury models and extended beyond those observed at walk–trot speeds only. By reproducing these behavioral changes in speed-dependent gait expression in a computational model of spinal locomotor circuitry, we identified potential principles of circuit reorganization underlying post-injury locomotor recovery following asymmetrical lateral hemisection and symmetrical contusion injuries at the low-thoracic level.

The model qualitatively recapitulated the *in vivo* locomotor characteristics observed after recovery from lateral hemisection when the ipsilesional descending drive and severed long propriospinal neurons were partially recovered, suggesting that the plastic changes responsible for recovery involved new and/or strengthened pathways that bypass the lesion to drive the hindlimb rhythm generators. In contrast, the model changes that effectively recapitulated locomotor recovery after a symmetrical midline contusion injury involved strengthening of the local left–right circuitry within each enlargement. These differences suggest that injury symmetry partly dictates the location and type of plasticity supporting recovery.

Despite these contrasts, the analyses converge on a shared requirement for recovered locomotion: reactivation of sublesional lumbar rhythm-generating circuits together with balanced lumbar commissural connectivity. Thus, asymmetric and symmetric injuries may rely on different pathways to achieve a common end point, but restored excitability of lumbar rhythm generators and effective commissural coordination consistently emerge as key determinants of post-injury gait expression.

Accordingly, clinical approaches — including neuromodulation and strategies to enhance regeneration or sprouting — may be more effective when aimed at restoring sublesional lumbar rhythm-generator and commissural function in a manner tailored to injury symmetry. Finally, we anticipate that further development of such computational approaches will eventually support the creation of individualized models to better characterize injury profiles and assist in development of therapeutic strategies.

Methods

Experimental data

In this modeling study, we used previously published experimental data on gait characteristics of overground locomotion within the full range of speeds in intact rats and following their recovery from either a mild-moderate contusion or a lateral hemisection spinal cord injury (Danner et al., 2023). In those studies, all experiments were performed in accordance with the Public Health Service Policy on Humane Care and Use of Laboratory Animals, and with the approval of the Institutional Animal Care and Use Committee (IACUC, protocol number 19644) at the University of Louisville. No new animals were used in the current study. Locomotion was assessed in a 3-m long Plexiglas tank where the walking surface was coated with sylgard to provide increased grip. A group of 17 Adult female Sprague-Dawley rats (225–250 g) was trained, using food treats, to traverse the length of the tank. Recordings were made using four high-speed (200 Hz) video cameras placed to capture the ventral view of the animal and analyzed using custom software. Then rats were divided into two groups and given either lateral hemisection (nine rats) or midline contusion injuries (eight rats; 12.5 g cm) at the T10 level of the spinal cord. The animals were allowed to recover for at least 4 weeks and were re-introduced to the long-tank set-up and trained again. At this time their locomotor recovery plateaued (contusion: 15.7 ± 1.9 , hemisection 18.3 ± 0.8 BBB Open Field Locomotor Scale). The post-recovery recordings were analyzed for each type of injury and compared to those for the intact animals. For more details, see Danner et al. (2023).

Modeling methods

Our model represents a network of interacting neural populations. Each population was simulated as an activity-based (firing rate) neuronal population model. In this description, the voltage variable V represents an average membrane potential of population and the output function $f(V)$ transverses V to the integrated population activity or normalized average firing rate at the corresponding voltage V (Ermentrout, 1994). This description was used in our previous modeling studies (Rubin et al., 2009; Danner et al., 2016, 2017; Ausborn et al., 2019; Latash et al., 2020; Rybak et al., 2024) and allows for inclusion of explicitly represented ionic currents, particularly, the leakage current I_L and the persistent (slowly inactivating) sodium current, I_{NaP} (Rubin et al., 2009). The latter current was proposed to be responsible for generating rhythmic activity in the spinal cord (Rybak et al., 2006; McCrea and Rybak, 2007; Tazerart et al., 2007, 2008; Brocard et al., 2013) and was included in description of flexor and extensor RG centers.

The variable V for flexor and extensor centers obeyed by the differential equation:

$$C \cdot \frac{dV}{dt} = -I_{NaP} - I_L - I_{SynE} - I_{SynI} - I_{Noise}. \quad (1)$$

All other populations did not include I_{NaP} , and in those populations, V was described as:

$$C \cdot \frac{dV}{dt} = -I_L - I_{SynE} - I_{SynI} - I_{Noise}, \quad (2)$$

where C is the membrane capacitance, I_{NaP} is the persistent sodium current, I_L is the leakage current,

I_{SynE} and I_{SynI} excitatory and inhibitory synaptic currents, respectively, and I_{Noise} a noisy current.

The leakage current was described as:

$$I_L = g_L \cdot (V - E_L), \quad (3)$$

where g_L is the leakage conductance and E_L represents the leakage reversal potential.

The persistent sodium current in the flexor and extensor half-centers was described as:

$$I_{NaP} = \bar{g}_{NaP} \cdot m(V) \cdot h(V) \cdot (V - E_{Na}), \quad (4)$$

where $\bar{g}_{\text{NaP}}$ is the I_{NaP} maximal conductance, E_{Na} is the sodium reversal potential, and $m(V)$ and $\bar{h}(V)$ are voltage-dependent activation and (slow) inactivation variables of I_{NaP} . $m(V)$ was instantaneous and its steady state was described as

$$m(V) = m_{\infty}(V) = \left\{1 + \exp\left[\frac{(V - V_{1/2,m})}{k_m}\right]\right\}^{-1}, \quad (5)$$

$$\tau_h(V) \cdot \frac{dh}{dt} = h_{\infty}(V) - h, \quad (6)$$

$$h_{\infty}(V) = \left\{1 + \exp\left[\frac{(V - V_{1/2,h})}{k_h}\right]\right\}^{-1}, \quad (7)$$

$$\tau_h(V) = \tau_0 + (\tau_{\max} - \tau_0) / \cosh\left[\frac{(V - V_{1/2,\tau})}{k_{\tau}}\right], \quad (8)$$

where $\bar{h}_{\infty}(V)$ is the inactivation steady state and $\tau_h(V)$ the inactivation time constant.

In Equations (5–8), $V_{1/2}$ and k represent half-voltage and slope of the corresponding variables (m , h , and τ); τ_0 and $\tau_{\max}$ are the baseline and maximum of inactivation time constant $\tau_{\bar{h}}$, respectively.

In Equation (1) and (2), excitatory and inhibitory synaptic currents (I_{SynE} and I_{SynI} , respectively) for population i were described as:

$$I_{\text{SynE},i} = g_{\text{SynE}} \cdot \{\sum_j [S(w_{ji}) \cdot f(V_j)] + D_{E,i}\} \cdot (V_i - E_{\text{SynE}}); \quad (9)$$

$$I_{\text{SynI},i} = g_{\text{SynI}} \cdot \{\sum_j [S(-w_{ji}) \cdot f(V_j)] + D_{I,i}\} \cdot (V_i - E_{\text{SynI}}), \quad (10)$$

where g_{SynE} and g_{SynI} are synaptic conductances and E_{SynE} and E_{SynI} are the reversal potentials of the excitatory and inhibitory synapses, respectively; w_{ji} is the synaptic weight from population j to population i ($w_{ji} > 0$ for excitatory connections and $w_{ji} < 0$ for inhibitory connections); function S is described as follows:

$$S(x) = \begin{cases} x, & \text{if } x \geq 0 \\ 0, & \text{if } x < 0 \end{cases}. \quad (11)$$

The output function $f(V)$ ($0 \leq f(V) \leq 1$) was defined as

$$f(V) = \begin{cases} 0, & \text{if } V < V_{\text{thr}} \\ \frac{V - V_{\text{thr}}}{V_{\max} - V_{\text{thr}}}, & \text{if } V_{\text{thr}} \leq V < V_{\max} \\ 1, & \text{if } V \geq V_{\max} \end{cases}. \quad (12)$$

Excitatory $D_{E,i}$ and inhibitory $D_{I,i}$ drives to population i were modeled as a linear function of the free parameter α :

$$D_{\{E,I\},i} = d_{\{E,I\},i} \cdot \alpha + b_{\{E,I\},i}, \quad (13)$$

where $d_{\{E,I\},i}$ is the slope and $b_{\{E,I\},i}$ is the intercept; the free parameter α represent the strength of the brainstem drive.

The noisy current $I_{\text{Noise},i}$ for each population i was described as an Ornstein–Uhlenbeck process

$$dI_{\text{Noise},i} = -\frac{I_{\text{Noise},i}}{\tau_{\text{Noise}}} dt + \sigma_{\text{Noise}} \cdot \sqrt{\frac{2}{\tau_{\text{Noise}}}} dW_i, \quad (14)$$

where τ_{Noise} is the time constant, σ_{Noise} the standard deviation of the noise, and W_i a Wiener process. It was numerically simulated using the finite difference formula

$$I_{\text{Noise},i}(t + \Delta t) = I_{\text{Noise},i}(t) - \frac{I_{\text{Noise},i}}{\tau_{\text{Noise}}} \Delta t + \sigma_{\text{Noise}} \cdot \sqrt{\frac{2\Delta t}{\tau_{\text{Noise}}}} \xi_i, \quad (15)$$

where is $\xi_i \sim \mathcal{N}(0,1)$ a normally-distributed random number with a variance of 1, and Δt the time step (1 ms).

Model parameters

The general neuron parameters were adapted from our previous models (Danner et al., 2016 [DOI](#), 2017 [DOI](#)) to provide the frequency range of locomotor activity characteristic for rats. The following values of parameters were used: $C = 10$ pF; $g_L = 4.5$ nS for RG centers and $g_L = 2.8$ nS for all other neurons; $\bar{g}_{NaP} = 4.5$ nS; $g_{synE} = g_{synI} = 10$ nS $E_L = -62.5$ mV for RG centers, $E_L = -60$ mV for all other populations; $E_{Na} = 50.0$ mV; $E_{EsynE} = -10$ mV; $E_{EsynI} = -75$ mV; $V_{thr} = -50$ mV; $V_{max} = 0$ mV; $V_{1/2,m} = -40.0$ mV; $k_m = -6$ mV; $V_{1/2,\square} = -45.0$ mV; $k_{\square} = 4$ mV; $\tau_{max} = 400$ ms; $\tau_0 = 150$ ms; $V_{1/2,\tau} = -35$ mV.

The following drive parameters were used for the intact model: $d_E = 0.0$ and $b_E = 0.1$ for the extensor centers; $d_E = 0.1$ and $b_E = 0.0$ for the flexor centers; $d_I = 0.75$ and $b_I = 0.0$ for homologous (within a girdle) $V0_D$ CINs in both cervical and lumbar compartments; $d_I = 1.5$ and $b_I = 0.0$ for diagonal $V0_D$ LPNs ($dV0_D$); $d_I = 0.25$ and $b_I = 0.0$ for the homologous cervical (fore) $V0_V$ CINs; $d_I = 0.15$ and $b_I = 0.0$ for the homologous lumbar (hind) $V0_V$ CINs. Connection weights for the intact model are listed in Table 1 [DOI](#).

To simulate rat behavior after recovery from right hemisection, drive to the ipsilesional lumbar flexor center was reduced by 10%, inhibitory drives to both cervical and lumbar $V0_V$ CINs were reduced by 50%, and connection weights of LPN connections affected by the injury were reduced to 40% of their pre-injury values (see Table 2 [DOI](#) for a comprehensive list of differences to the pre-injury case).

To simulate rat behavior after recovery from contusion, all weights of LPN connection were reduced by 95%. In addition, the increasing inhibitory drives to lumbar $V0_D$ and $V0_V$ CINs were eliminated and replaced by constant drives (see Table 3 [DOI](#) for a comprehensive list of differences to the pre-injury case).

Computer simulations

The set of differential equations were solved using the odeint (Ahnert et al., 2011 [DOI](#)) implementation of the Cash-Karp 5th-order Runge Kutta method (Cash and Karp, 1990 [DOI](#)) of the boost C++ library (version 1.88.0). The custom C++ code was compiled (with clang-1700.0.13.3 for MAC OS 15.3.1) as a Python package. Python 3.11.5 (Python Software Foundation, Wilmington, Delaware, USA) was used to run simulations and further analyze the results. The neural network simulation package is available at <https://github.com/SimonDanner/CPGNetworkSimulator> [DOI](#).

Data analysis

Activities of the flexor and extensor centers were used to assess the model behavior. Each rhythm generator was considered as being in flexion when output $f(V)$ of its flexor center was greater or equal to 0.1 otherwise it was considered as being in extension. The midextension time was defined as the halfway between onset and offset of extension. The locomotor period (step) was defined as the duration between two consecutive midextension time points; the frequency as the reciprocal of the period; extension and flexion phase durations as the duration between on- and offset of extension and flexion, respectively. Normalized phase differences were calculated as the delay between the midextension time points of a pair of rhythm generators divided by the locomotor period. Six normalized phase differences were calculated: hind left–right (right hind–left hind), fore left–right (right fore–left fore), left homolateral (left fore–left hind), right homolateral (right fore–right hind), and two diagonal (right fore–left hind and left fore–right hind).

Gait classification

Gaits for each step cycle were operationally defined based on phase differences, similarly to the method used in Danner et al. (2023) [DOI](#). Each step was represented as a point Φ in the phase space consisting of the three (non-normalized) phase differences with the left hind RG as a reference. Then the distance to all idealized one-, two-, three-, and four-beat gaits (T_g ; Table 4 [DOI](#)), was calculated

Table 1. Connection weights in intact model

Source	Target (w_{ij})
Within cervical and lumbar circuits	
RG-F	i-InF (0.4), i-V0 _D (0.7), i-V2a (1)
RG-E	i-InE (0.4), i-V3-E (0.35), i-Sh2 (0.5)
IniF	i-RG-E (-1)
IniE	i-RG-F (-0.1)
V2a	i-V0 _V (1)
V0 _V	c-Ini (0.6)
V0 _D	c-RG-F (-0.07)
V3-E	c-RG-E (0.02)
Within cervical circuits	
RG-F	i-dLPNi (0.7), i-dV0 _D (0.5), i-dV2a (0.5)
Ini	i-RG-F (-0.0375)
dV2a	i-dV0 _V (0.9)
Within lumbar circuits	
RG-F	i-V3-F (0.4), i-aV3 (0.3)
Ini	i-RG-F (-0.075)
V3-F	c-RG-F (0.03)
V3-E	c-InE1 (1)
InE1	c-RG-E (-0.045)
Between cervical and lumbar circuits	
dSh2	ih-RG-F (0.005)
aSh2	if-RG-F (0.04)
dLPNi	ih-RG-F (-0.01)
dV0 _D	ch-RG-F (-0.075)
dV0 _V	ch-RG-F (0.02)
aV3	cf-RG-F (0.065)

i-, ipsilateral; c-, contralateral; f-, fore; h-, hind.

Table 2. Hemisection: differences to pre-injury model

Connection weights	
Source	Target ($w_{\text{intact}} \rightarrow w_{\text{hemisection}}$)
hl-aV3	fl-RG-F (0.065 $\rightarrow$ 0.026)
hr-aSh2	fr-RG-F (0.04 $\rightarrow$ 0.016)
fr-dSh2	hr-RG-F (0.005 $\rightarrow$ 0.002)
fr-dLPNi	hr-RG-F (-0.01 $\rightarrow$ -0.004)
fl-dV0 _D	hl-RG-F (-0.075 $\rightarrow$ -0.03)
fl-dV0 _V	hl-RG-F (0.02 $\rightarrow$ 0.008)
Drive parameters	
Target	$d_{\{E,I\},\text{intact}} \rightarrow d_{\{E,I\},\text{hemisection}}$
rh-RG-F	$d_E = 0.1 \rightarrow d_E = 0.009$
h-V0 _V	$d_I = 0.15 \rightarrow d_I = 0.075$
f-V0 _V	$d_I = 0.25 \rightarrow d_I = 0.125$

f-, fore; h-, hind; l-, left; r-, right.

Connection weights	
Source	Target ($w_{\text{intact}} \rightarrow w_{\text{contusion}}$)
h-aV3	cf-RG-F (0.065 $\rightarrow$ 0.00325)
h-aSh2	if-RG-F (0.04 $\rightarrow$ 0.002)
f-dSh2	ih-RG-F (0.005 $\rightarrow$ 0.00025)
f-dLPNi	ih-RG-F (-0.01 $\rightarrow$ -0.0005)
f-dV0 _D	ch-RG-F (-0.075 $\rightarrow$ -0.00375)
f-dV0 _V	ch-RG-F (0.02 $\rightarrow$ 0.001)
Drive parameters	
Target	$d/b_{\{E,I\},\text{intact}} \rightarrow d/b_{\{E,I\},\text{contusion}}$
h-V0 _D	$d_I = 0.15 \rightarrow d_I = 0.0$; $b_I = 0.0 \rightarrow b_I = 0.2$
h-V0 _V	$d_I = 0.25 \rightarrow d_I = 0.0$; $b_I = 0.0 \rightarrow b_I = 0.03$

i-, ipsilateral; c-, contralateral; f-, fore; h-, hind.

Table 3. Contusion: differences to pre-injury model

$$d(\Phi, \Psi_g) = \sqrt{\sum_{j \in \{LR, HL, DIAG\}} \arg[e^{i(\phi_j - \psi_j)}]^2} \quad (16)$$

and the step cycle was assigned to belong to the gait $\mathbf{T}_g$ to which it was closest. The duty factor was not considered for gait classification. Gait transitions were analyzed on a step-by-step basis (Danner et al., 2023 [↗](#)).

Analysis of model performance

To model step-to-step variability and gait transitions comparable with locomotor bouts of rats in the long-tank (Danner et al., 2023 [↗](#)), simulations with increased noisy currents ($\sigma_{\text{Noise}} = 1.1 \text{ pA}$) were performed in all three conditions: intact, post-hemisection, and post-contusion. In these simulations, α was gradually ramped up from 0.55 to 1.05 for the intact condition, or to 1.0 for hemisection and contusion conditions, and then ramped back down to 0.55. Each phase of the ramp lasted 8 seconds, and the entire ramp cycle was repeated 200 times. Variability of phase differences was quantified as the deviation of each phase difference from its circular moving average (cutoff frequency 0.125 step cycles) averaged across ramp cycles (Danner et al., 2023 [↗](#)).

To identify stable solutions of the model, bifurcation diagrams (Danner et al., 2017 [↗](#)) were built for the six normalized phase differences. For this purpose, α was incrementally increased from 0.075 to 1.1, and then decreased back to 0.075, using a step size of 0.001. At each step, the simulation was run for 10 s, then if the normalized circular standard deviation of the five last step cycles of each phase difference was smaller than 0.005, the final state was used as the initial condition for the next step, otherwise, the same procedure was repeated. If the normalized standard deviation was not met by 100 s of simulation, the model was considered not stable at this specific α -value. Otherwise, the mean phase differences were regarded as the stable solutions. Initial conditions were randomized, and multiple runs were evaluated. To ensure that the model does not remain on an unstable trajectory, these simulations were performed with a weak noisy current ($\sigma_{\text{Noise}} = 5 \text{ fA}$).

Sensitivity analysis

To identify which connectivity changes most strongly influence post-injury gait, we performed a local sensitivity analysis of the injured (contusion and hemisection) models. For each model, we quantified how parameter variation altered locomotor behavior relative to the baseline parameter set. Simulations with noise were performed for each parameter perturbation, as described above but with 25 ramp cycles.

As a summary outcome, we computed the Earth Mover's Distance (EMD; Rubner et al., 1998 [↗](#)) between the baseline simulation and each perturbed simulation in a 7-dimensional feature space defined by the six interlimb phase differences and locomotor frequency. EMD measures the minimal “work” required to transform one multivariate pattern into another and thus provides a single, interpretable metric of overall locomotor pattern disruption. Distances were computed as the sum of circular (torus) distances for the six interlimb phase differences combined with the Euclidean distance for frequency.

Model parameters were perturbed multiplicatively on the logarithmic scale $p' = p \cdot e^x$, $x \sim \text{Uniform}[-\ln r, \ln r]$, where p is the baseline parameter value, and p' the perturbed parameter value. We used $r = 1.25$, so that the scale factors ranged over $[1/r, r] = [0.8, 1.25]$. This approach ensured that parameters remained strictly positive and that their increases and decreases were treated as symmetric percentage changes rather than absolute shifts.

For each analysis, we generated a Sobol quasi-Monte Carlo training sample of 2^{13} perturbations, ran the simulation, and computed the corresponding EMD values. In addition, we generated and evaluated an independent Sobol validation set of 2^9 samples with a different random seed. This dataset was reserved exclusively for surrogate validation.

We trained an XGBoost gradient-boosted tree model (Friedman, 2001 [↗](#); Chen and Guestrin, 2016 [↗](#)) as a surrogate (Storlie et al., 2009 [↗](#)) to approximate the mapping of the perturbed parameters to the EMD to the baseline model. Surrogate accuracy was then assessed on the

	Normalized phase differences			Gait
	LR	HL	diag.	
One-beat	0	0	0	Pronk
Two-beat	1/2	1/2	0	Trot
	0	1/2	1/2	Bound
	0	2/3	2/3	Bound
	1/2	0	1/2	Pace
Three-beat	0	1/3	2/3	Half-bound
	0	2/3	1/3	Half-bound
	2/3	1/3	0	Canter
	1/3	1/3	2/3	Canter
	1/3	2/3	0	Other
	2/3	2/3	1/3	Other
	1/3	2/3	2/3	Other
	2/3	1/3	1/3	Other
	1/3	0	2/3	Other
	2/3	0	1/3	Other
	1/3	2/3	1/3	Other
	2/3	1/3	2/3	Other
Four-beat	3/4	1/4	2/4	Rotary gallop
	1/4	3/4	2/4	Rotary gallop
	3/4	2/4	1/4	Transverse gallop
	1/4	2/4	3/4	Transverse gallop
	2/4	1/4	3/4	Lateral-sequence
	2/4	3/4	1/4	Diagonal-sequence

LR: hind left-right normalized phase difference, HL: homolateral normalized phase difference; diag.: diagonal normalized phase difference. Adapted from Danner et al. (2023).

Table 4. Idealized gaits

independent validation set: validation R^2 was within 5% of that of training R^2 and Spearman rank correlation between simulated and surrogate predictions exceeded 0.95.

Sobol sensitivity indices were then estimated using the surrogate model (Sobol', 2001 [link](#); Storlie et al., 2009 [link](#)). We used Saltelli sampling (Saltelli et al., 2010 [link](#)), a variance-based Monte Carlo method that generates structured resampling matrices to efficiently compute both first-order and total-order indices. Saltelli sampling was performed with $N \cdot (k+2)$ samples, with k being the number of parameters and $N=10^{14}$. Using the surrogate, we estimated first-order and total-order Sobol indices via quasi-Monte Carlo sampling. Indices are reported with 95% confidence intervals. All indices should be interpreted as variance contributions within the local neighborhood based on the perturbation band [80%,125%], not as global sensitivities across the entire parameter space. Note that reported total-order indices include all interaction terms involving the feature and thus their sum can be greater than 1.

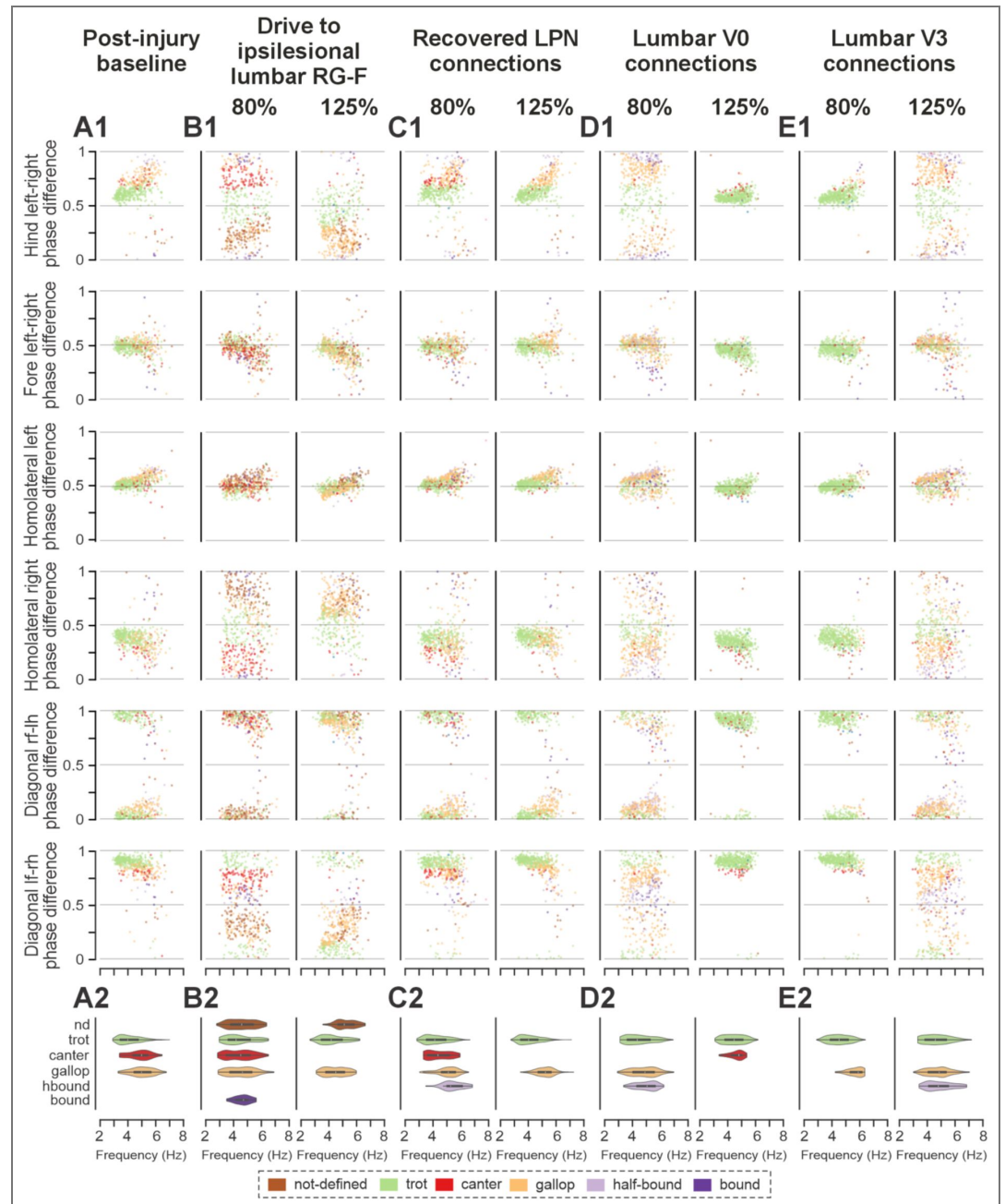


Figure 13—figure supplement 1. Effects of parameter variation on model output in the hemisection model.

Shown are scatter plots of normalized phase-difference vs. frequency distributions (A1–E1) and corresponding gait–frequency distributions (A2–E2) for the post-injury baseline model (A) and for four parameters varied individually to the lower (80%) and upper (125%) bounds of the sensitivity analysis range: drive to the ipsilesional lumbar flexor rhythm generator center (RG-F) (B), recovered long propriospinal neuron (LPN) connections (C), lumbar V0 commissural interneuron (CIN) connections (D), and lumbar V3 CIN connections (E). All other parameters were held at their baseline values. l-, left; r-, right; -f, fore; -h, hind RG.

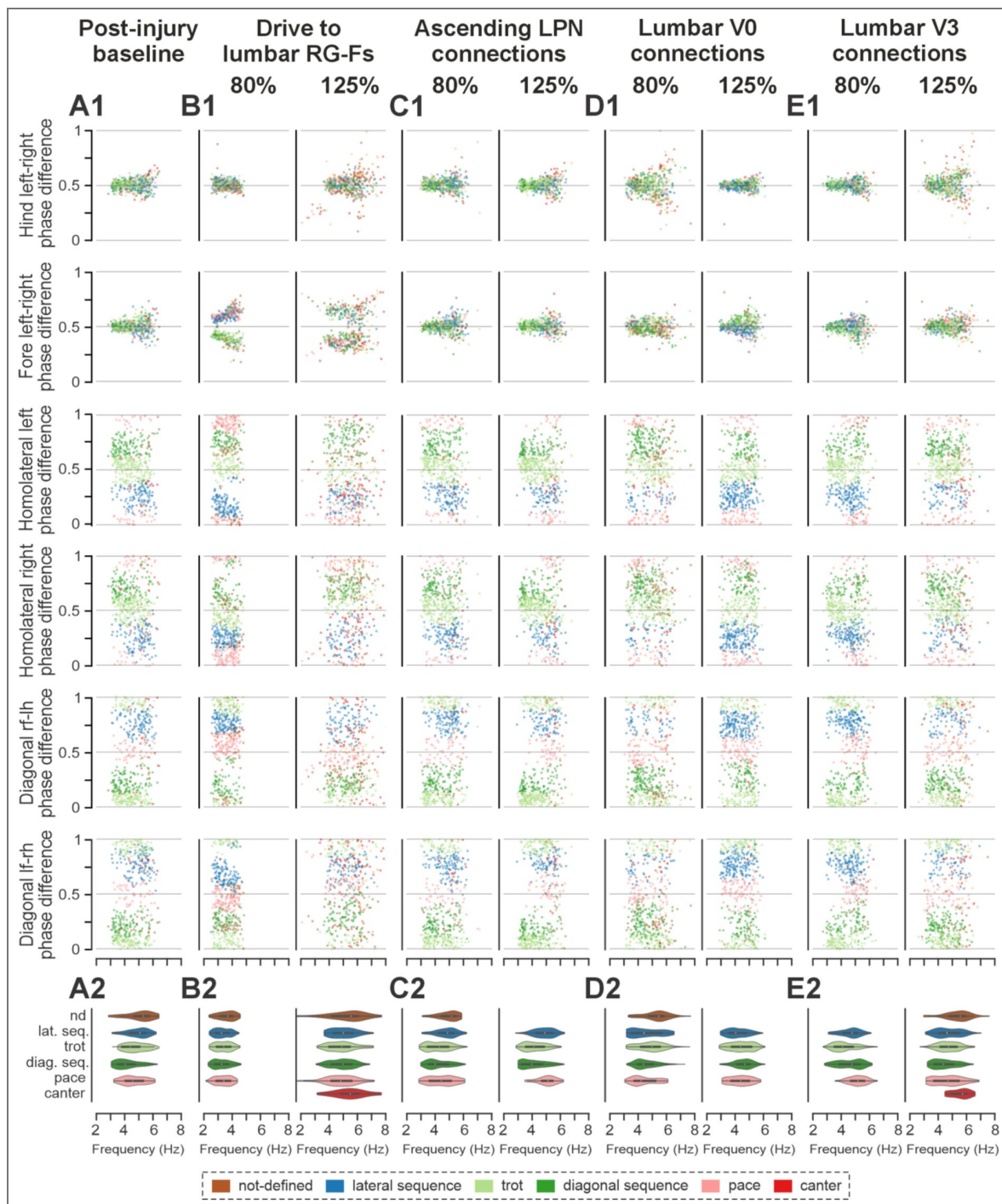


Figure 13—figure supplement 2. Effects of parameter variation on model output in the contusion model.

Shown are scatter plots of normalized phase-difference vs. frequency distributions (A1–E1; random sample of 500 locomotor cycles) and corresponding gait-frequency distributions (A2–E2) for the post-injury baseline model (A) and for four parameters varied individually to the lower (80%) and upper (125%) bounds of the sensitivity analysis range: drive to the lumbar flexor rhythm generator centers (RG-Fs) (B), ascending long propriospinal neuron (LPN) connections (C), lumbar V0 commissural interneuron (CIN) connections (D), and lumbar V3 CIN connections (E). All other parameters were held at their baseline values. l-, left; r-, right; -f, fore; -h, hind RG; nd, not-defined (gait); lat. seq., lateral-sequence; diag. seq., diagonal-sequence.

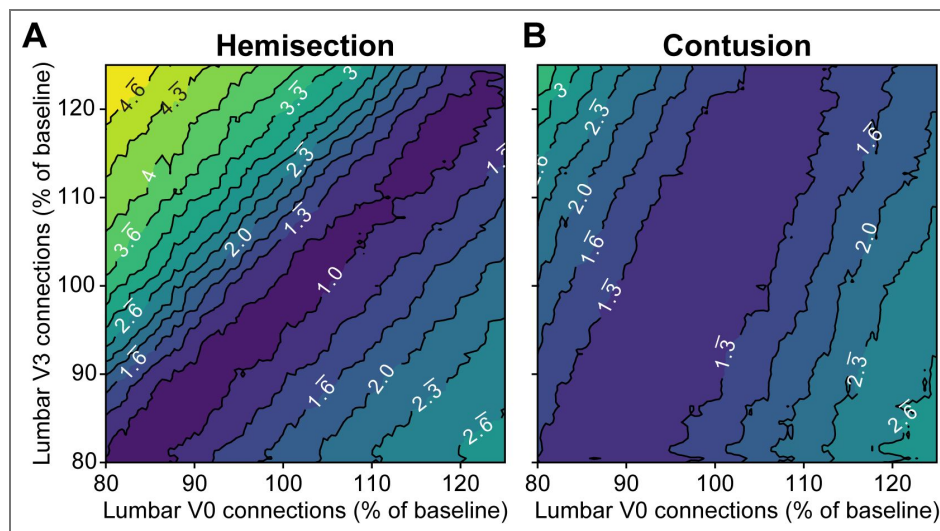


Figure 13—figure supplement 3. Predicted Earth Mover's Distance to baseline as a function of lumbar V0/V3 commissural interneuron connectivity.

Contour plots of surrogate-predicted Earth Mover's Distance as a function of lumbar V0 and V3 commissural interneuron connection strengths (80–125% of baseline; same range as used for the sensitivity analysis) in the hemisection (A) and contusion (B) models, with all other parameters held at their baseline values.

Data Availability

Model configuration files, Python code for running simulations, analyzing results, and processing experimental data, as well as scripts to reproduce all simulations and generate all figures presented in the paper, are available at <https://github.com/dannerlab/rat-sci-locomotion-model>. The neural network simulation package used in this study is available at <https://github.com/SimonDanner/CPGNetworkSimulator>. The experimental data reproduced from Danner et al. (2023) can be found at <https://github.com/dannerlab/rat-sci-locomotion>.

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References

- Abbinanti MD**, Harris-Warrick RM (2012) Serotonin modulates multiple calcium current subtypes in commissural interneurons of the neonatal mouse. *Journal of Neurophysiology* **107**:2212-2219 <https://doi.org/10.1152/jn.00768.2011>
- Abbinanti MD**, Zhong G, Harris-Warrick RM (2012) Postnatal emergence of serotonin-induced plateau potentials in commissural interneurons of the mouse spinal cord. *Journal of Neurophysiology* **108**:2191-2202 <https://doi.org/10.1152/jn.00336.2012>
- Ahnert K**, Mulansky M, Simos TE, Psihoyios G, Ch Tsitouras, Anastassi Z (2011) Odeint – Solving Ordinary Differential Equations in C++. *AIP Conf Proc* **1389**:1586-1589 <https://doi.org/10.1063/1.3637934>
- Akay T**, Tourtellotte WG, Arber S, Jessell TM (2014) Degradation of mouse locomotor pattern in the absence of proprioceptive sensory feedback. *Proc Natl Acad Sci* **111**:16877-16882 <https://doi.org/10.1073/pnas.1419045111>

- Aoi S, Kondo T, Hayashi N, Yanagihara D, Aoki S, Yamaura H, Ogihara N, Funato T, Tomita N, Senda K, *et al.* (2013) Contributions of phase resetting and interlimb coordination to the adaptive control of hindlimb obstacle avoidance during locomotion in rats: A simulation study. *Biol Cybern* **107**:201-216 <https://doi.org/10.1007/s00422-013-0546-6>
- Ausborn J, Shevtsova NA, Caggiano V, Danner SM, Rybak IA (2019) Computational modeling of brainstem circuits controlling locomotor frequency and gait. *eLife* **8**:e43587 <https://doi.org/10.7554/eLife.43587>
- Ausborn J, Shevtsova NA, Danner SM (2021) Computational Modeling of Spinal Locomotor Circuitry in the Age of Molecular Genetics. *Ijms* **22**:6835 <https://doi.org/10.3390/ijms22136835>
- Ballermann M, Fouad K (2006) Spontaneous locomotor recovery in spinal cord injured rats is accompanied by anatomical plasticity of reticulospinal fibers. *European Journal of Neuroscience* **23**:1988-1996 <https://doi.org/10.1111/j.1460-9568.2006.04726.x>
- Ballion B, Morin D, Viala D (2001) Forelimb locomotor generators and quadrupedal locomotion in the neonatal rat. *Eur J Neurosci* **14**:1727-1738 <https://doi.org/10.1046/j.0953-816X.2001.01794.x>
- Barbeau H, Rossignol S (1987) Recovery of locomotion after chronic spinalization in the adult cat. *Brain Res* **412**:84-95
- Barriere G, Leblond H, Provencher J, Rossignol S (2008) Prominent Role of the Spinal Central Pattern Generator in the Recovery of Locomotion after Partial Spinal Cord Injuries. *Journal of Neuroscience* **28**:3976-3987 <https://doi.org/10.1523/JNEUROSCI.5692-07.2008>
- Basso DM, Beattie MS, Bresnahan JC (1996) Graded histological and locomotor outcomes after spinal cord contusion using the NYU weight-drop device versus transection. *Exp Neurol* **139**:244-256 <https://doi.org/10.1006/exnr.1996.0098>
- Bellardita C, Kiehn O (2015) Phenotypic characterization of speed-associated gait changes in mice reveals modular organization of locomotor networks. *Curr Biol* **25**:1426-1436 <https://doi.org/10.1016/j.cub.2015.04.005>
- Bertram JE (2016) *Understanding mammalian locomotion: concepts and applications* Hoboken, New Jersey: John Wiley & Sons.
- Brocard F, Shevtsova NA, Bouhadjane M, Tazerart S, Heinemann U, Rybak IA, Vinay L (2013) Activity-dependent changes in extracellular Ca²⁺ and K⁺ reveal pacemakers in the spinal locomotor-related network. *Neuron* **77**:1047-1054 <https://doi.org/10.1016/j.neuron.2013.01.026>
- Brown BL, Anil N, States G, Whittemore SR, Magnuson DSK (2024) Long ascending propriospinal neurons are heterogeneous and subject to spinal cord injury induced anatomic plasticity. *Experimental Neurology* **373**:114631 <https://doi.org/10.1016/j.expneurol.2023.114631>
- Burns AS, Ditunno JF (2001) Establishing Prognosis and Maximizing Functional Outcomes After Spinal Cord Injury Rehabilitation Management. *Spine* **26**:137-145
- Butt SJB, Kiehn O (2003) Functional identification of interneurons responsible for left-right coordination of hindlimbs in mammals. *Neuron* **38**:953-963 [https://doi.org/10.1016/S0896-6273\(03\)00353-2](https://doi.org/10.1016/S0896-6273(03)00353-2)
- Caggiano V, Leiras R, Goni-Erro H, Masini D, Bellardita C, Bouvier J, Caldeira V, Fisone G, Kiehn O (2018) Midbrain circuits that set locomotor speed and gait selection. *Nature* **553**:455-460 <https://doi.org/10.1038/nature25448>
- Capelli P, Pivetta C, Soledad Esposito M, Arber S (2017) Locomotor speed control circuits in the caudal brainstem. *Nature* **551**:373-377 <https://doi.org/10.1038/nature24064>
- Cash JR, Karp AH (1990) A variable order Runge-Kutta method for initial value problems with rapidly varying right-hand sides. *ACM Trans Math Softw* **16**:201-222 <https://doi.org/10.1145/79505.79507>
- Chen T, Guestrin C. (2016) XGBoost: A Scalable Tree Boosting System. In: Proceedings of the 22nd ACM SIGKDD International Conference on Knowledge Discovery and Data Mining. San Francisco, United States. pp. 785-794 <https://doi.org/10.1145/2939672.2939785>

- Cohen AH, Gans C (1975) Muscle activity in rat locomotion: Movement analysis and electromyography of the flexors and extensors of the elbow. *J Morphol* **146**:177-196 <https://doi.org/10.1002/jmor.1051460202>
- Cohen-Adad J, Martinez M, Delivet-Mongrain H, Rossignol S (2014) Recovery of locomotion after partial spinal cord lesions in cats: assessment using behavioral, electrophysiological and imaging techniques. *Acta Neurobiol Exp* **74**:142-157
- Courtine G, Song B, Roy RR, Zhong H, Herrmann JE, Ao Y, Qi J, Edgerton VR, Sofroniew MV (2008) Recovery of supraspinal control of stepping via indirect propriospinal relay connections after spinal cord injury. *Nat Med* **14**:69-74 <https://doi.org/10.1038/nm1682>
- Crone SA, Zhong G, Harris-Warrick R, Sharma K (2009) In mice lacking V2a interneurons, gait depends on speed of locomotion. *J Neurosci* **29**:7098-7109 <https://doi.org/10.1523/JNEUROSCI.1206-09.2009>
- Danner SM, Hofstoetter US, Freundl B, Binder H, Mayr W, Rattay F, Minassian K (2015) Human spinal locomotor control is based on flexibly organized burst generators. *Brain* **138**:577-588 <https://doi.org/10.1093/brain/awu372>
- Danner SM, Shepard CT, Hainline C, Shevtsova NA, Rybak IA, Magnuson DSK (2023) Spinal control of locomotion before and after spinal cord injury. *Experimental Neurology* **368**:114496 <https://doi.org/10.1016/j.expneurol.2023.114496>
- Danner SM, Shevtsova NA, Frigon A, Rybak IA (2017) Computational modeling of spinal circuits controlling limb coordination and gaits in quadrupeds. *eLife* **6**:e31050 <https://doi.org/10.7554/eLife.31050>
- Danner SM, Wilshin SD, Shevtsova NA, Rybak IA (2016) Central control of interlimb coordination and speed-dependent gait expression in quadrupeds. *J Physiol* **594**:6947-6967 <https://doi.org/10.1113/jp272787>
- Danner SM, Zhang H, Shevtsova NA, Borowska-Fielding J, Deska-Gauthier D, Rybak IA, Zhang Y (2019) Spinal V3 Interneurons and Left-Right Coordination in Mammalian Locomotion. *Front Cell Neurosci* **13**:516 <https://doi.org/10.3389/fncel.2019.00516>
- Dutton RC, Carstens MI, Antognini JF, Carstens E (2006) Long ascending propriospinal projections from lumbosacral to upper cervical spinal cord in the rat. *Brain Res* **1119**:76-85 <https://doi.org/10.1016/j.brainres.2006.08.063>
- Edgley SA, Jankowska E, Krutki P, Hammar I (2003) Both dorsal horn and lamina VIII interneurons contribute to crossed reflexes from feline group II muscle afferents. *J Physiol* **552**:961-974 <https://doi.org/10.1113/jphysiol.2003.048009>
- Edinger L (1896) *Vorlesungen über den Bau der nervösen Centralorgane des Menschen und der Thiere* (5th ed) Leipzig, Deutschland: Verlag von F.C.W. Vogel.
- Engmann AK, Bizzozzero F, Schneider MP, Pfyffer D, Imobersteg S, Schneider R, Hofer A-S, Wieckhorst M, Schwab ME (2020) The Gigantocellular Reticular Nucleus Plays a Significant Role in Locomotor Recovery after Incomplete Spinal Cord Injury. *J Neurosci* **40**:8292-8305 <https://doi.org/10.1523/JNEUROSCI.0474-20.2020>
- Ermentrout B (1994) Reduction of Conductance-Based Models with Slow Synapses to Neural Nets. *Neural Comput* **6**:679-695 <https://doi.org/10.1162/neco.1994.6.4.679>
- Filli L, Engmann AK, Zorner B, Weinmann O, Moraitis T, Gullo M, Kasper H, Schneider R, Schwab ME (2014) Bridging the Gap: A Reticulo-Propriospinal Detour Bypassing an Incomplete Spinal Cord Injury. *J Neurosci* **34**:13399-13410 <https://doi.org/10.1523/JNEUROSCI.0701-14.2014>
- Filli L, Schwab M (2015) Structural and functional reorganization of propriospinal connections promotes functional recovery after spinal cord injury. *Neural Regen Res* **10**:509 <https://doi.org/10.4103/1673-5374.155425>
- Flynn JR, Conn VL, Boyle KA, Hughes DI, Watanabe M, Velasquez T, Goulding MD, Callister RJ, Graham BA (2017) Anatomical and Molecular Properties of Long Descending Propriospinal Neurons in Mice. *Front Neuroanat* **11** <https://doi.org/10.3389/fnana.2017.00005>

- Forssberg H, Grillner S, Halbertsma J, Rossignol S (1980) The locomotion of the low spinal cat. II. Interlimb coordination. *Acta Physiol Scand* **108**:283-295 <https://doi.org/10.1111/j.1748-1716.1980.tb06534.x>
- Fouad K, Tse A (2008) Adaptive changes in the injured spinal cord and their role in promoting functional recovery. *Neurol Res* **30**:17-27 <https://doi.org/10.1179/016164107X251781>
- Friedman JH (2001) Greedy function approximation: A gradient boosting machine. *Ann Statist* **29** <https://doi.org/10.1214/aos/1013203451>
- Frigon A (2017) The neural control of interlimb coordination during mammalian locomotion. *J Neurophysiol* **117**:2224-2241 <https://doi.org/10.1152/jn.00978.2016>
- Frigon A, Gossard J (2012) Asymmetric control of cycle period by the spinal locomotor rhythm generator in the adult cat. *J Physiol* **587**:4617-4627 <https://doi.org/10.1113/jphysiol.2009.176669>
- Grillner S. (1981) Control of locomotion in bipeds, tetrapods, and fish. *Comprehensive Physiology* 1179-1236
- Halbertsma JM (1983) The stride cycle of the cat: the modelling of locomotion by computerized analysis of automatic recordings. *Acta Physiol Scand Suppl* **521**:1-75
- Hildebrand M (1989) The Quadrupedal Gaits of Vertebrates. *Bioscience* **39**:766-775 <https://doi.org/10.2307/1311182>
- Hsu L-J, Bertho M, Kiehn O (2023) Deconstructing the modular organization and real-time dynamics of mammalian spinal locomotor networks. *Nat Commun* **14**:873 <https://doi.org/10.1038/s41467-023-36587-w>
- Jankowska E (2008) Spinal interneuronal networks in the cat: Elementary components. *Brain Research Reviews* **57**:46-55 <https://doi.org/10.1016/j.brainresrev.2007.06.022>
- Jankowska E, Bannatyne BA, Stecina K, Hammar I, Cabaj A, Maxwell DJ (2009) Commissural interneurons with input from group I and II muscle afferents in feline lumbar segments: neurotransmitters, projections and target cells. *J Physiol* **587**:401-418 <https://doi.org/10.1113/jphysiol.2008.159236>
- Jordan LM, Liu J, Hedlund PB, Akay T, Pearson KG (2008) Descending command systems for the initiation of locomotion in mammals. *Brain Res Rev* **57**:183-191 <https://doi.org/10.1016/j.brainresrev.2007.07.019>
- Josset N, Roussel M, Lemieux M, Lafrance-Zoubga D, Rastqar A, Bretzner F (2018) Distinct contributions of mesencephalic locomotor region nuclei to locomotor control in the freely behaving mouse. *Current Biology* **28**:884-901 <https://doi.org/10.1016/j.cub.2018.02.007>
- Juvin L, Le Gal J-P, Simmers J, Morin D. (2012) Cervicolumbar coordination in mammalian quadrupedal locomotion: role of spinal thoracic circuitry and limb sensory inputs. *J Neurosci* **32**:953-965 <https://doi.org/10.1523/JNEUROSCI.4640-11.2012>
- Kato M (1990) Chronically isolated lumbar half spinal cord generates locomotor activities in the ipsilateral hindlimb of the cat. *Neurosci Res* **9**:22-34 [https://doi.org/10.1016/0168-0102\(90\)90042-D](https://doi.org/10.1016/0168-0102(90)90042-D)
- Kiehn O (2006) Locomotor circuits in the mammalian spinal cord. *Annu Rev Neurosci* **29**:279-306 <https://doi.org/10.1146/annurev.neuro.29.051605.112910>
- Kiehn O (2016) Decoding the organization of spinal circuits that control locomotion. *Nat Rev Neurosci* **17**:224-238 <https://doi.org/10.1038/nrn.2016.9>
- Kim Y, Aoi S, Fujiki S, Danner SM, Markin SN, Ausborn J, Rybak IA, Yanagihara D, Senda K, Tsuchiya K (2022) Contribution of Afferent Feedback to Adaptive Hindlimb Walking in Cats: A Neuromusculoskeletal Modeling Study. *Front Bioeng Biotechnol* **10**:825149 <https://doi.org/10.3389/fbioe.2022.825149>
- Kjaerulff O, Kiehn O (1997) Crossed rhythmic synaptic input to motoneurons during selective activation of the contralateral spinal locomotor network. *J Neurosci* **17**:9433-9447

- Kuerzi J, Brown EH, Shum-Siu A, Siu A, Burke D, Morehouse J, Smith RR, Magnuson DSK (2010) Task-specificity vs. ceiling effect: step-training in shallow water after spinal cord injury. *Exp Neurol* **224**:178-187 <https://doi.org/10.1016/j.expneurol.2010.03.008>
- Laflamme OD, Markin SN, Deska-Gauthier D, Banks R, Zhang Y, Danner SM, Akay T (2023) Distinct roles of spinal commissural interneurons in transmission of contralateral sensory information. *Current Biology* **33**:3452-3464.e4. <https://doi.org/10.1016/j.cub.2023.07.014>
- Latah EM, Lecomte CG, Danner SM, Frigon A, Rybak IA, Molkov YI (2020) On the Organization of the Locomotor CPG: Insights From Split-Belt Locomotion and Mathematical Modeling. *Front Neurosci* **14**:598888 <https://doi.org/10.3389/fnins.2020.598888>
- Lemieux M, Josset N, Roussel M, Couraud S, Bretzner F (2016) Speed-dependent modulation of the locomotor behavior in adult mice reveals attractor and transitional gaits. *Front Neurosci* **10**:42 <https://doi.org/10.3389/fnins.2016.00042>
- Lemieux M, Karimi N, Bretzner F (2024) Functional plasticity of glutamatergic neurons of medullary reticular nuclei after spinal cord injury in mice. *Nat Commun* **15**:1542 <https://doi.org/10.1038/s41467-024-45300-4>
- Lloyd DPC, McIntyre AK (1948) Analysis of forelimb-hindlimb reflex activity in acutely decapitate cats. *J Neurophysiol* **11**:455-470
- Lovely RG, Gregor RJ, Roy RR, Edgerton VR (1986) Effects of training on the recovery of full-weight-bearing stepping in the adult spinal cat. *Exp Neurol* **92**:421-435
- McCrea DA, Rybak IA (2008) Organization of mammalian locomotor rhythm and pattern generation. *Brain Res Rev* **57**:134-146 <https://doi.org/10.1016/j.brainresrev.2007.08.006>
- McCrea DA, Rybak IA (2007) Modeling the mammalian locomotor CPG: insights from mistakes and perturbations. *Prog Brain Res* **165**:235-253 [https://doi.org/10.1016/S0079-6123\(06\)65015-2](https://doi.org/10.1016/S0079-6123(06)65015-2)
- Molkov YI, Yu G, Ausborn J, Bouvier J, Danner SM, Rybak IA (2024) Sensory feedback and central neuronal interactions in mouse locomotion. *R Soc Open Sci* **11**:240207
- Nathan PW, Smith MC (1959) Fasciculi proprii of the spinal cord in man. *Brain* **82**:610-668
- Nishikawa K, Biewener AA, Aerts P, Ahn AN, Chiel HJ, Daley MA, Daniel TL, Full RJ, Hale ME, Hedrick TL, et al. (2007) Neuromechanics: An integrative approach for understanding motor control. *Integr Comp Biol* **47**:16-54 <https://doi.org/10.1093/icb/icm024>
- Orlovsky GN, Deliagina TG, Grillner S (1999) *Neuronal control of locomotion: from mollusc to man* New York: Oxford University Press.
- Perrier J-F, Cotel F (2015) Serotonergic modulation of spinal motor control. *Current Opinion in Neurobiology* **33**:1-7 <https://doi.org/10.1016/j.conb.2014.12.008>
- Pocratsky AM, Shepard CT, Morehouse JR, Burke DA, Riegler AS, Hardin JT, Beare JE, Hainline C, States GJ, Brown BL, et al. (2020) Long ascending propriospinal neurons provide flexible, context-specific control of interlimb coordination. *eLife* **9**:e53565 <https://doi.org/10.7554/eLife.53565>
- Raineteau O, Schwab ME (2001) Plasticity of motor systems after incomplete spinal cord injury. *Nat Rev Neurosci* **2**:263-273 <https://doi.org/10.1038/35067570>
- Reed WR, Shum-Siu A, Onifer SM, Magnuson DSK (2006) Inter-enlargement pathways in the ventrolateral funiculus of the adult rat spinal cord. *Neuroscience* **142**:1195-1207 <https://doi.org/10.1016/j.neuroscience.2006.07.017>
- Rosenzweig ES, Courtine G, Jindrich DL, Brock JH, Ferguson AR, Strand SC, Nout YS, Roy RR, Miller DM, Beattie MS, et al. (2010) Extensive spontaneous plasticity of corticospinal projections after primate spinal cord injury. *Nat Neurosci* **13**:1505-10 <https://doi.org/10.1038/nn.2691>
- Rossignol S, Dubuc R, Gossard J-P (2006) Dynamic sensorimotor interactions in locomotion. *Physiol Rev* **86**:89-154 <https://doi.org/10.1152/physrev.00028.2005>
- Rubin JE, Shevtsova NA, Ermentrout GB, Smith JC, Rybak IA (2009) Multiple rhythmic states in a model of the respiratory central pattern generator. *J Neurophysiol* **101**:2146-2165 <https://doi.org/10.1152/jn.90958.2008>

- Rubner Y, Tomasi C, Guibas LJ (1998) A metric for distributions with applications to image databases. In: Sixth International Conference on Computer Vision (IEEE Cat. No.98CH36271). Bombay, India. pp. 59-66 <https://doi.org/10.1109/ICCV.1998.710701>
- Ruder L, Takeoka A, Arber S (2016) Long-Distance Descending Spinal Neurons Ensure Quadrupedal Locomotor Stability. *Neuron* **92**:1063-1078 <https://doi.org/10.1016/j.neuron.2016.10.032>
- Rybak IA, Dougherty KJ, Shevtsova NA (2015) Organization of the mammalian locomotor cpg: Review of computational model and circuit architectures based on genetically identified spinal interneurons. *eNeuro* **2**:ENEURO.0069-15.2015 <https://doi.org/10.1523/ENEURO.0069-15.2015>
- Rybak IA, Shevtsova NA, Audet J, Yassine S, Markin SN, Prilutsky BI, Frigon A (2025) Operation of spinal sensorimotor circuits controlling phase durations during tied-belt and split-belt locomotion after a lateral thoracic hemisection. *eLife* **13**:RP103504 <https://doi.org/10.7554/eLife.103504.3>
- Rybak IA, Shevtsova NA, Lafreniere-Roula M, McCrea DA (2006) Modelling spinal circuitry involved in locomotor pattern generation: insights from deletions during fictive locomotion. *J Physiol* **577**:617-639 <https://doi.org/10.1113/jphysiol.2006.118703>
- Rybak IA, Shevtsova NA, Markin SN, Prilutsky BI, Frigon A (2024) Operation regimes of spinal circuits controlling locomotion and the role of supraspinal drives and sensory feedback. *eLife* **13**:RP98841 <https://doi.org/10.7554/eLife.98841.3>
- Saltelli A, Annoni P, Azzini I, Campolongo F, Ratto M, Tarantola S (2010) Variance based sensitivity analysis of model output. Design and estimator for the total sensitivity index. *Computer Physics Communications* **181**:259-270 <https://doi.org/10.1016/j.cpc.2009.09.018>
- Shepard CT, Brown BL, Van Rijswijck MA, Zalla RM, Burke DA, Morehouse JR, Riegler AS, Whittemore SR, Magnuson DS. (2023) Silencing long-descending inter-enlargement propriospinal neurons improves hindlimb stepping after contusive spinal cord injuries. *eLife* **12**:e82944 <https://doi.org/10.7554/eLife.82944>
- Shepard CT, Pocratsky AM, Brown BL, Van Rijswijck MA, Zalla RM, Burke DA, Morehouse JR, Riegler AS, Whittemore SR, Magnuson DS. (2021) Silencing long ascending propriospinal neurons after spinal cord injury improves hindlimb stepping in the adult rat. *eLife* **10**:e70058 <https://doi.org/10.7554/eLife.70058>
- Shevtsova NA, Talpalar AE, Markin SN, Harris-Warrick RM, Kiehn O, Rybak IA (2015) Organization of left-right coordination of neuronal activity in the mammalian spinal cord: Insights from computational modelling. *J Physiol* **593**:2403-2426 <https://doi.org/10.1113/jp270121>
- Sobol' IM. (2001) Global sensitivity indices for nonlinear mathematical models and their Monte Carlo estimates. *Mathematics and Computers in Simulation* **55**:271-280 [https://doi.org/10.1016/S0378-4754\(00\)00270-6](https://doi.org/10.1016/S0378-4754(00)00270-6)
- Storlie CB, Swiler LP, Helton JC, Sallaberry CJ (2009) Implementation and evaluation of nonparametric regression procedures for sensitivity analysis of computationally demanding models. *Reliability Engineering & System Safety* **94**:1735-1763 <https://doi.org/10.1016/j.ress.2009.05.007>
- Takeoka A, Arber S (2019) Functional local proprioceptive feedback circuits initiate and maintain locomotor recovery after spinal cord injury. *Cell Reports* **27**:71-85 <https://doi.org/10.1016/j.celrep.2019.03.010>
- Takeoka A, Vollenweider I, Courtine G, Arber S (2014) Muscle spindle feedback directs locomotor recovery and circuit reorganization after spinal cord injury. *Cell* **159**:1626-1639 <https://doi.org/10.1016/j.cell.2014.11.019>
- Talpalar AE, Bouvier J, Borgius L, Fortin G, Pierani A, Kiehn O (2013) Dual-mode operation of neuronal networks involved in left-right alternation. *Nature* **500**:85-88 <https://doi.org/10.1038/nature12286>
- Tazerart S, Viemari J-C, Darbon P, Vinay L, Brocard F (2007) Contribution of persistent sodium current to locomotor pattern generation in neonatal rats. *J Neurophysiol* **98**:613-628 <https://doi.org/10.1152/jn.00316.2007>

Tazerart S, Vinay L, Brocard F (2008) The persistent sodium current generates pacemaker activities in the central pattern generator for locomotion and regulates the locomotor rhythm. *J Neurosci* **28**:8577-8589 <https://doi.org/10.1523/JNEUROSCI.1437-08.2008>

Zhang H, Shevtsova NA, Deska-Gauthier D, Mackay C, Dougherty KJ, Danner SM, Zhang Y, Rybak IA (2022) The role of V3 neurons in speed-dependent interlimb coordination during locomotion in mice. *eLife* **11**:e73424 <https://doi.org/10.7554/eLife.73424>

Zörner B, Bachmann LC, Filli L, Kapitza S, Gullo M, Bolliger M, Starkey ML, Röthlisberger M, Gonzenbach RR, Schwab ME (2014) Chasing central nervous system plasticity: the brainstem's contribution to locomotor recovery in rats with spinal cord injury. *Brain* **137**:1716-1732 <https://doi.org/10.1093/brain/awu078>

Peer reviews

Reviewer #1 (Public review):

Summary:

This is a rigorous data-driven modeling study extending the authors' previous model of spinal locomotor central pattern generator (CPG) circuits developed for the mouse spinal cord and adapted here to the rat to explore potential circuit-level changes underlying altered speed-dependent gaits due to asymmetric (lateral) thoracic spinal hemisection and symmetric midline contusion. The model reproduces key features of the rat speed-dependent gait-related experimental data before injury and after recovery from these two different thoracic spinal cord injuries and suggests injury-specific mechanisms of circuit reorganization underlying functional recovery. There is much interest in the mechanisms of locomotor behavior recovery after spinal cord injury, and data-driven behaviorally relevant circuit modeling is an important approach. This study represents an important advance of the authors' previous experimental and modeling work on locomotor circuitry and in the motor control field.

Strengths:

- (1) The authors use an advanced computational model of spinal locomotor circuitry to investigate potential reorganization of neural connectivity underlying locomotor control following recovery from symmetrical midline thoracic contusion and asymmetrical (lateral) hemisection injuries, based on an extensive dataset for the rat model of spinal cord injury.
- (2) The rat dataset used is from an in vivo experimental paradigm involving challenging animals to perform overground locomotion across the full range of speeds before and after the two distinct spinal cord injury models, enabling the authors to more completely reveal injury-specific deficits in speed-dependent interlimb coordination and locomotor gaits.
- (3) The model reproduces the rat gait-related experimental data before injury and after recovery from these two different thoracic spinal cord injuries, which exhibit roughly comparable functional recovery, and suggests injury-specific, compensatory mechanisms of circuit reorganization underlying recovery.
- (4) The model simulations suggest that recovery after lateral hemisection mechanistically involves partial functional restoration of descending drive and long propriospinal pathways, whereas recovery following midline contusion relies on reorganization of sublesional lumbar circuitry combined with altered descending control of cervical networks.
- (5) These observations suggest that symmetrical (contusion) and asymmetrical (lateral hemisection) injuries induce distinct types of plasticity in different spinal cord regions, suggesting that injury symmetry partly dictates the location and type of neural plasticity supporting recovery.

(6) The authors suggest therapeutic strategies may be more effective by targeting specific circuits according to injury symmetry.

Weaknesses:

(1) The recovery mechanisms implemented in the model involve circuit connectivity/connection weights adjustment based on assumptions about the structures involved and compensatory responses to the injury. As the authors acknowledge, other factors affecting locomotor patterns and compensation, such as somatosensory afferent feedback, neurochemical modulator influences, and limb/body biomechanics, are not considered in the model. The authors have now more adequately discussed the limitations of the modeling and associated implications for functional interpretation.

Comments on revisions:

The authors have substantially improved the manuscript by including model parameter sensitivity analyses and by more adequately discussing the limitations of the modeling and associated implications for functional interpretation.

<https://doi.org/10.7554/eLife.107480.2.sa3>

Reviewer #2 (Public review):

Summary:

In this paper, the authors present a detailed computational model and experimental data concerning over-ground locomotion in rats before and after recovery from spinal cord injury. They are able to manually tune the parameters of this physiologically based, detailed model to reproduce many aspects of the observed animals' locomotion in the naive case and in two distinct injury cases.

Strengths:

The strengths are that the model is driven to closely match clean experimental data, and the model itself has detailed correspondence to proposed anatomical reality. As such this makes the model more readily applicable to future experimental work. It can make useful suggestions. The model reproduces a large number of conditions, across frequencies, and with model structure changed by injury and recovery. The model is extensive and is driven by known structures, has links to genetic identities, and has been validated extensively across a number of experiments and manipulations over the years. It models a system of critical importance to the field, and the tight coupling to experimental data is a real strength.

Weaknesses:

A downside is that scientifically, here, the only question tackled is one of sufficiency. With manual tuning of parameters in a way that matches what the field believes/knows from experimental work, the detailed model can reproduce the experimental findings. One of the benefits of computational models is that the counter-factual can be tested to provide evidence against alternate hypotheses. That isn't really done here. I'm pretty sure there are competing theories of what happens during recovery from a hemi-section injury and contusion injury. The model could be used to make predictions for some alternate hypothesis, supporting or rejecting theories of recovery. This may be part of future plans. Here, the focus is on showing that the model is capable of reproducing the experimental results at all, for any set of parameters, however tuned.

Comments on revisions:

The authors have addressed my prior concerns and clearly discuss the sufficiency of the model, and strengthen the discussion with interesting findings for the role of propriospinal and commissural interneuronal pathways. This is a very nice contribution.

<https://doi.org/10.7554/eLife.107480.2.sa2>

Reviewer #3 (Public review):

Summary:

This study describes a computational model of the rat spinal locomotor circuit and how it could be reconfigured after lateral hemisection or contusion injuries to replicate gaits observed experimentally.

The model suggests the emergence of detour circuits after lateral hemisection whereas after a midline contusion, the model suggests plasticity of left-right and sensory inputs below the injury.

Strengths:

The model accurately models many known connections within and between forelimb and hindlimb spinal locomotor circuits.

The simulation results mirror closely gait parameters observed experimentally. Many gait parameters were studied as well as variability in these parameters in intact versus injured conditions.

A sensitivity analysis provides some sense of the relative importance of the various modified connectivity after injury in setting the changes in gait seen after the two types of injuries

Overall, the authors achieved their aims and the model provides solid support for the changes in connectivity after the two types of injuries modelled. This work emphasizes specific changes in connectivity after lateral hemisection or after contusion that could be investigated experimentally. The model is available to be used by the public and could be a tool used to investigate the relative importance of various highlighted or undiscovered changes in connectivity that could underlie the recovery of locomotor function in spinalized rats.

Comments on revisions:

The authors addressed the comments made by the reviewers. The sensitivity analysis adds insights to the manuscript

<https://doi.org/10.7554/eLife.107480.2.sa1>

Author response:

The following is the authors' response to the original reviews.

Reviewer #1 (Public review):

Summary:

This is a rigorous data-driven modeling study, extending the authors' previous model of spinal locomotor central pattern generator (CPG) circuits developed for the mouse spinal cord and adapted here to the rat to explore potential circuit-level changes underlying altered speeddependent gaits, due to asymmetric (lateral) thoracic spinal hemisection

and symmetric midline contusion. The model reproduces key features of the rat speed-dependent gait-related experimental data before injury and after recovery from these two different thoracic spinal cord injuries and suggests injury-specific mechanisms of circuit reorganization underlying functional recovery. There is much interest in the mechanisms of locomotor behavior recovery after spinal cord injury, and data-driven behaviorally relevant circuit modeling is an important approach. This study represents an important advance in the authors' previous experimental and modeling work on locomotor circuitry and in the motor control field.

Strengths:

(1) The authors use an advanced computational model of spinal locomotor circuitry to investigate potential reorganization of neural connectivity underlying locomotor control following recovery from symmetrical midline thoracic contusion and asymmetrical (lateral) hemisection injuries, based on an extensive dataset for the rat model of spinal cord injury.

(2) The rat dataset used is from an in vivo experimental paradigm involving challenging animals to perform overground locomotion across the full range of speeds before and after the two distinct spinal cord injury models, enabling the authors to more completely reveal injury-specific deficits in speed-dependent interlimb coordination and locomotor gaits.

(3) The model reproduces the rat gait-related experimental data before injury and after recovery from these two different thoracic spinal cord injuries, which exhibit roughly comparable functional recovery, and suggests injury-specific, compensatory mechanisms of circuit reorganization underlying recovery.

(4) The model simulations suggest that recovery after lateral hemisection mechanistically involves partial functional restoration of descending drive and long propriospinal pathways. In contrast, recovery following midline contusion relies on reorganization of sublesional lumbar circuitry combined with altered descending control of cervical networks.

(5) These observations suggest that symmetrical (contusion) and asymmetrical (lateral hemisection) injuries induce distinct types of plasticity in different spinal cord regions, suggesting that injury symmetry partly dictates the location and type of neural plasticity supporting recovery.

(6) The authors suggest that therapeutic strategies may be more effective by targeting specific circuits according to injury symmetry.

Weaknesses:

The recovery mechanisms implemented in the model involve circuit connectivity/connection weights adjustment based on assumptions about the structures involved and compensatory responses to the injury. As the authors acknowledge, other factors affecting locomotor patterns and compensation, such as somatosensory afferent feedback, neurochemical modulator influences, and limb/body biomechanics, are not considered in the model.

We appreciate the positive review and critical comments. We added a dedicate limitation and future direction section (see response recommendations below). Further, we also performed a sensitivity analysis: while the model still relies on a set of hypothesized connectivity changes, this analysis quantifies how robust our conclusions are to these parameter choices and indicates which pathways most strongly affect the recovered locomotor pattern.

Reviewer #1 (Recommendations for the authors):

The authors have used an advanced model of rodent spinal locomotor CPG circuits, adapted to the rat spinal cord, which remarkably reproduces the key features of the rat speed-dependent gait-related experimental data before injury and after recovery from the two different thoracic spinal cord injuries studied. Importantly, they have exploited the extensive dataset for the in vivo rat spinal cord injury model involving overground locomotion across the full range of speeds before and after the two distinct spinal cord injuries, enabling the authors to more completely reveal injury-specific deficits in speed-dependent interlimb coordination and locomotor gaits. The paper is well-written and well-illustrated.

(1) My only general suggestion is that the authors include a section that succinctly summarizes the limitations of the modeling and points to elaborations of the model and experimental data required for future studies. Some important caveats are dispersed throughout the Discussion, but a more consolidated section would be useful.

We added a dedicated Limitations and future directions section (page XX) that consolidates shortcomings and broadly outlines potential next steps in terms of modeling and experimental data. Specifically, we highlight the issue of lack of afferent feedback connections in the model, lack of consideration of biomechanic mechanisms, and restriction of the model to beneficial plasticity. To resolve these issues, we need neuromechanical models (integration of the neural circuits with a model of the musculoskeletal system), experimental data validating our predictions and data to constrain future models to be able to distinguish between beneficial and maladaptive plasticity.

(2) Please correct the Figure 11 legend title to indicate recovery after contusion (not hemisection).

Done. Thanks for noticing.

Reviewer #2 (Public review):

Summary:

In this paper, the authors present a detailed computational model and experimental data concerning overground locomotion in rats before and after recovery from spinal cord injury. They are able to manually tune the parameters of this physiologically based, detailed model to reproduce many aspects of the observed animals locomotion in the naive case and in two distinct injury cases.

Strengths:

The strengths are that the model is driven to closely match clean experimental data, and the model itself has detailed correspondence to proposed anatomical reality. As such, this makes the model more readily applicable to future experimental work. It can make useful suggestions. The model reproduces a large number of conditions across frequencies, and with the model structure changed by injury and recovery. The model is extensive and is driven by known structures, with links to genetic identities, and has been extensively validated across multiple experiments and manipulations over the years. It models a system of critical importance to the field, and the tight coupling to experimental data is a real strength.

Weaknesses:

A downside is that, scientifically, here, the only question tackled is one of sufficiency. By manually tuning parameters in a manner that aligns with the field's understanding from

experimental work, the detailed model can accurately reproduce the experimental findings. One of the benefits of computational models is that the counterfactual can be tested to provide evidence against alternative hypotheses. That isn't really done here. I'm fairly certain that there are competing theories regarding what happens during recovery from a hemi-section injury and a contusion injury. The model could be used to make predictions for some alternative hypotheses, supporting or rejecting theories of recovery. This may be part of future plans. Here, the focus is on showing that the model is capable of reproducing the experimental results at all, for any set of parameters, however tuned.

We agree with the reviewer that the present study focuses on sufficiency, and we now explicitly acknowledge this in the revised limitations section. We also added sensitivity analysis (for details see response to reviewer 3) that provides an initial assessment of robustness to the assumed connectivity changes. We note that the model reproduces a broad set of experimentally observed features across the full range of locomotor frequencies (including loss and emergence of specific gaits, reduced maximum stepping frequency, and altered variability of interlimb phase differences) using only a small set of hypothesized circuit reorganizations that have been experimentally observed but previously only correlated with recovery. Our results therefore suggest that this limited set of changes is indeed sufficient to account for the complex pattern of recovered locomotor behavior.

Finally, although exploring alternative solutions is of interest, we believe such efforts will be most informative once afferent feedback is incorporated, which we see as the logical next step in our studies.

Reviewer #2 (Recommendations for the authors):

The paper could be strengthened with some more scientific interpretation and future directions. What are some novel predictions that can be made with the model, now that it has shown sufficiency here, that could guide future experimental work? Does it contradict in any way theories of CPG structure or neuronal plastic recovery?

The sensitivity analysis that we performed in response to reviewer 3's suggestion expanded our interpretation/conclusions by showing that, although injury symmetry (contusion vs. lateral hemisection) influences *which* pathways reorganize, recovered locomotion across injury type depends most strongly on restored activation of lumbar rhythm-generating and strengths of lumbar commissural circuits.

Interestingly, this sensitivity analysis also showed that variations of strengths of long propriospinal pathways (ascending, descending, spared, injured-and-recovered) have a much smaller, almost negligible effect, when compared to variations of drive to lumbar rhythm generators or lumbar commissural interneuron connection weights in the same range (see Fig 13, 13-supplement 1 and 2). This is in accordance with our initial model suggestions that after contusion LPN connections weight had to be lowered to values substantially lower than what was expected by the severity of the injury. Which is also corroborated by our anatomical findings that in parallel to recovery from contusion, the number of synaptic connections by LAPNs to the cervical enlargement were reduced, and that silencing of LPNs post-contusion improves locomotion. These surprising findings have been extensively discussed in the discussion section.

Together, these findings suggest that experimental characterization of reorganization of the lumbar circuitry with a specific focus on commissural interneurons and inputs to the lumbar circuitry that could restore activation of sublesional lumbar rhythm generators is a crucial next step for understanding post-injury plasticity and recovery of locomotor function. This is now clearly discussed.

Finally, we note that a key contribution of this work is that the model demonstrates a plausible mechanistic link between specific circuit reorganizations and recovered locomotor

function, a relationship previously supported mainly by correlative evidence.

Reviewer #3 (Public review):

Summary:

This study describes a computational model of the rat spinal locomotor circuit and how it could be reconfigured after lateral hemisection or contusion injuries to replicate gaits observed experimentally.

The model suggests the emergence of detour circuits after lateral hemisection, whereas after a midline contusion, the model suggests plasticity of left-right and sensory inputs below the injury.

Strengths:

The model accurately models many known connections within and between forelimb and hindlimb spinal locomotor circuits.

The simulation results mirror closely gait parameters observed experimentally. Many gait parameters were studied, as well as variability in these parameters in intact versus injured conditions.

Weaknesses:

The study could provide some sense of the relative importance of the various modified connectivities after injury in setting the changes in gait seen after the two types of injuries.

We performed a local sensitivity analysis of the hemisection and contusion models to identify which connectivity changes most strongly influence post-injury locomotor behavior. Key parameters (descending drive to sublesional rhythm generators and the strength of selected commissural and propriospinal pathways) were perturbed within 80–125% of their baseline values, and for each perturbation we quantified changes in model output using the Earth Mover's Distance between baseline and perturbed simulations in a 7-dimensional space (six interlimb phase differences plus locomotor frequency). We then trained a surrogate model and computed Sobol first- and total-order sensitivity indices, which quantify how much each parameter and its interactions contribute to variability in this distance measure. This analysis showed that, across both injuries, variations in drive to sublesional lumbar rhythm generators and in lumbar V0/V3 commissural connectivity have the largest impact on recovered gait expression, whereas other pathways had comparatively minor effects within the tested range.

The sensitivity analysis further refined our conclusions by showing that, although injury symmetry (contusion vs. lateral hemisection) influences *which* pathways reorganize, effective recovery in both cases depends on re-engaging lumbar rhythm-generating and commissural circuits, highlighting these networks as key therapeutic targets.

Overall, the authors achieved their aims, and the model provides solid support for the changes in connectivity after the two types of injuries were modelled. This work emphasizes specific changes in connectivity after lateral hemisection or after contusion that could be investigated experimentally. The model is available for public use and could serve as a tool to analyze the relative importance of various highlighted or previously undiscovered changes in connectivity that may underlie the recovery of locomotor function in spinalized rats.

Reviewer #3 (Recommendations for the authors):

(1) It would be useful to study the sensitivity of the injured models to small changes in the connectivity changes to determine which ones play a greater role in the gait after injury.

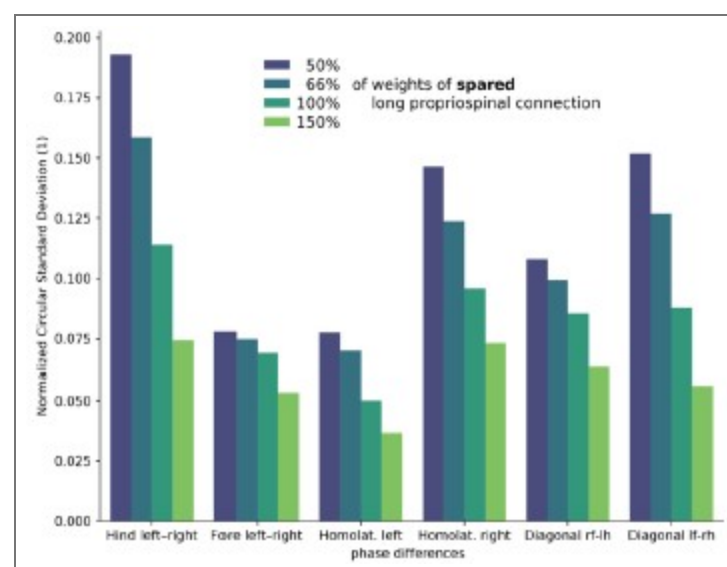
See response above on the added sensitivity analysis.

(2) Was there any tissue analysis from the original experiments with the contusion experiments, as contusion experiments can be variable, so it would be good to know the level of variability in the injuries?

Unfortunately, we were unable to complete tissue analysis of the injury epicenters for these animals because the tissue was not handled appropriately for histology. However, in the past, comparable animals with T10 12.5g-cm contusion injuries delivered by the NYU (MASCIS) Impactor had variability of up to ~30% of the mean (spared white matter, e.g. see Smith et al., 2006). It is also worth noting that spared white matter at the epicenter, at least in our hands, is generally well-correlated with BBB overground locomotor scale scores.

(3) There is more variability in phase difference in rats than model in the lateral hemisection. Is there any way to figure out which of the connectivity changes is most responsible for that variability?

We agree that the variability of phase differences after lateral hemisection is larger in rats than in the model. One possible contributor to this discrepancy is the strength of spared long propriospinal neuron (LPN) pathways, which we kept fixed at pre-injury levels in the model. As an exploratory analysis, we varied the weights of these spared LPN connections and quantified the circular standard deviation of the phase differences (Author response image 1). Decreasing spared LPN weights increased the variability of all phase differences. This suggests that plasticity of spared LPNs (potentially reducing their effective connectivity and partly compensating for the asymmetry introduced by the lesion) could contribute to the higher variability seen in vivo. However, because these results remain speculative, we chose to include them in this response only and not in the main manuscript.



Author response image 1. Variability of phase differences as a function of spared long propriospinal neuron connection weights (hemisection model).

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