GABA facilitates spike propagation at branch points of sensory axons in the spinal cord

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Abstract

Movement and posture depend on sensory feedback that is regulated by specialized GABAergic neurons (GAD2⁺) that form axo-axonic contacts onto myelinated proprioceptive sensory axons and are thought to be inhibitory. However, we report here that activating GAD2⁺ neurons, directly with optogenetics or indirectly by cutaneous stimulation, facilitates sensory feedback to motoneurons in awake rodents and humans. GABA_A receptors and GAD2⁺ contacts adjacent to nodes of Ranvier at branch points of sensory axons cause this facilitation, preventing spike propagation failure that is otherwise common without GABA. GABA_A receptors are generally lacking from axon terminals (unlike GABA_B) and do not inhibit transmitter release onto motoneurons, disproving the long-standing assumption that GABA_A receptors cause presynaptic inhibition. GABAergic innervation of nodes near branch points allows individual branches to function autonomously, with GAD2⁺ neurons regulating which branches conduct, adding a computational layer to the neuronal networks generating movement and likely generalizing to other CNS axons.

Main

The ease with which animals move defies the complexity of the underlying neuronal circuits, which include corticospinal tracts (CSTs) that coordinate skilled movement, spinal interneurons that form central patterns generators (CGPs) for walking, and motoneurons that ultimately drive the muscles¹. Sensory feedback ensures the final precision of such motor acts, with proprioceptive feedback to motoneurons producing a major part of the muscle activity in routine movement and posture²⁻⁴, without which severe ataxia occurs⁵. Proprioceptive sensory feedback is regulated by specialized GABAergic neurons (GAD2⁺; abbreviated GABA_{axo} neurons) that form axo-axonic connections onto the sensory axon terminals⁶⁻⁸. These neurons are thought to produce presynaptic inhibition of sensory feedback to motoneurons⁹⁻¹¹ and possibly limit inappropriate sensory feedback^{3,4,7}. However, during movement the CST, CPG and even sensory neurons all augment GABA_{axo} neuron activity¹¹⁻¹⁶ right at a time when sensory feedback is known to be increased to ensure precision and postural stability²⁻⁴, raising the question of whether GABA_{axo} neurons have a yet undescribed excitatory action.

The long-standing view that GABAergic neurons produce presynaptic inhibition of proprioceptive sensory axon terminals in adult mammals actually lacks direct evidence, largely because of the difficulty in recording from these small terminals and the technical limitations of previously employed

reduced spinal cord preparations (immature)^{7,17} or anesthetized animals, since anesthetics themselves modulate GABA receptors^{9,18}. Thus, in this paper we used optogenetic approaches to directly target GABA_{axo} neurons in awake animals and in isolated whole adult spinal cord preparations. Surprisingly, we found that optogenetically activating these GABAergic neurons markedly facilitates sensory axon transmission to motoneurons, throwing into doubt presynaptic inhibition.

GABA_A receptors on sensory axon terminals have long been assumed to cause presynaptic inhibition, based on indirect evidence^{11,18}, though their mode of action is rather counterintuitive. Sensory axons, like many other axons, have high intracellular chloride concentrations, leading to an outward chloride ion flow through activated GABA_A receptors^{11,19,20}. Thus, GABA_A receptors cause a depolarization of sensory axons (primary afferent depolarization, PAD)^{11,16,21-23}. PAD and associated GABA_A receptors have variously been theorized to cause presynaptic inhibition by depolarization-dependent inactivation or shunting of sodium currents at the sensory axon terminals^{22,24}. However, we do not even know if terminals of large myelinated proprioceptive sensory axons express GABA_A receptors at all, despite their demonstrated innervation by GABA_{axo} neurons⁶. These terminals appear to lack the α5 subunit of extrasynaptic GABA_A receptors¹⁶, but this leaves open the possibility that they express synaptic GABA_A or GABA_B receptors. We thus examined this question here and unexpectedly found that synaptic GABA_A receptors are generally not at these axon terminals, but are instead near sodium channels (Nav) at the nodes of Ranvier throughout the myelinated regions of the axon, together with innervation by GABA_{axo} neurons. What then is the function of such nodal GABA_A receptors?

An unexplored possibility is that the depolarizing action of nodal GABAA receptors (and GABA_{axo} neurons) aids sodium spike propagation between axon nodes. This has not previously been considered, as spikes are thought to securely propagate from node to node, at least in the orthodromic direction¹⁶. Myelinated proprioceptive axons branch extensively in the spinal cord¹⁶(Fig. 1a) and each branch point poses a theoretical risk for spike propagation failure^{25,26}. However, branch points are always located near nodes (Nav)¹⁶, likely to minimize this failure. Nevertheless, indirect evidence has suggested that propagation failure can occur²⁷⁻³⁰. Thus, in the present study we sought direct evidence of nodal spike failure at branch points and examined whether nodal GABA and PAD facilitates afferent conduction by preventing this failure. We already know that PAD lowers the threshold for initiating axon spikes by extracellular stimulation³¹, and even initiates spikes¹⁶, but do not know whether it aids normal spike propagation. We found that spike propagation depends so heavily on nodal GABA that blocking GABA action makes the majority of proprioceptive sensory axons fail to propagate spikes to motoneurons, and

thus GABA provides a powerful mechanism to turn on specific nodes and branches to regulate sensory feedback.

Results

Nodal GABA_A and terminal GABA_B receptors.

To visualize how GABA regulates axons we examined the distribution of GABAA receptor subunit immunolabelling on nodes and terminals in large myelinated proprioceptive sensory axons, including α 5 subunits that are extrasynaptic, α 1 and α 2 subunits that are mostly synaptic, and ubiquitous γ 2 subunits³². We labelled sensory axons with neurobiotin injections in the rat spinal cord (Fig. 1) or fluorescent reporters in VGLUT1^{Cre/+} mice (Extended Data Fig. 1), and reconstructed them in 3D. GABA_A receptors containing $\alpha 5$, $\alpha 1$, $\alpha 2$ and $\gamma 2$ subunits were expressed on these axons, especially near sodium channels (< 6 µm away; Fig. 1d-h, Extended Data Fig. 1). Specifically, GABAA receptors were on large myelinated 1st and 2nd order branches in the dorsal and ventral horn (Figs. 1d-e,h; Extended Data Fig. 1) near their nodes (identified by large sodium channel clusters, paranodal Caspr, and an axonal taper; < 10 µm away; Figs. 1c,h,j), and on short unmyelinated terminal branches in the dorsal horn (3rd order; Figs. 1b,g), the latter near nodes on 2nd order branches (< 100 µm away). In contrast, GABAA receptors were mostly absent from the long unmyelinated terminal branches contacting motoneurons in the ventral horn (3rd order; Figs. 1b,f,g; Extended Data Fig. 1a-e), which also generally lacked sodium channels¹⁶. This left GABA_A receptors on average far from the terminal boutons contacting motoneurons ($\sim 500 \, \mu m$; Fig. 1h) relative to the axon space constant ($\lambda_s \sim 90 \, \mu m$), the majority in dorsal and intermediate laminae. Nodes were widely spaced, as were branch points (~50 µm separation, Fig. 1i), but branch points were always near nodes (100%, within 7 µm; Nay; Fig. 1c-e,i) and associated GABAA receptors. While nodes sometimes occurred without branching (49%), the majority of nodes expressing GABAA receptors were at branch points (Fig. 1j), implying an importance for branch point GABA. In contrast, GABA_B receptors were found mostly on terminal branches in the ventral horn where boutons had dense receptor expression (Figs. 1f-h), and not usually on larger myelinated ventral or dorsal branches (not at nodes; Figs. 1e,g,j; Extended Data Fig. 1f).

Propagation failure in dorsal horn axon branches.

Considering that GABAA receptors are expressed in large myelinated branches of proprioceptive axons, we next directly recorded from of these branches in the dorsal horn of rat and mouse spinal cords (Figs.

2 and 3) to examine whether spike propagation depends on these receptors. When we stimulated the dorsal root (DR) containing the axon branch, an all-or-nothing spike was recorded in many branches (Figs. 2b, 3d) at the latency of the fastest afferent volley that arrived at the spinal cord (group I afferents; EC in Fig. 2b). However, in other axon branches this spike did not occur (~20%), but at the same latency there was a small all-or-nothing residual spike (*failure potential, FP*). This FP was indicative of a spike activating a distant node, but failing to propagate further to the recording site, leaving only its passively attenuated potential, with smaller FPs reflecting more distal failure points in the spinal cord (Figs. 2c-g, 3e-f). Failure never occurred in the DR itself (Fig. 2f). The failing branches with FPs were otherwise indistinguishable from non-failing axon branches, exhibiting full spikes (> 60 mV) with current injection pulses (Fig. 2cii, g), and low conductances and resting potentials (~ -65 mV, Fig. 2h), ruling out penetration injury. With high repetitive DR stimulation rates all branches (100%) exhibited propagation failure and an associated FP (Fig. 2e-g), again with the FP implying that the spike is reliably initiated in the DR, but incompletely propagates within the spinal cord.

Axon spike failure was voltage dependent: in branches with failing spikes (FPs) depolarizations that brought the axon closer to threshold enabled full DR-evoked spikes (via current injection, Fig. 2ci; or spontaneous depolarization Fig. 2d). Also, in branches without spike failure at rest (secure spikes) a steady hyperpolarizing current induced spike failure (FP), with more branches failing with increasing hyperpolarization (Extended Data Fig. 2). With increasing hyperpolarization, nodes failed progressively more distal to the electrode, causing abrupt drops in the overall spike amplitude with each failure and a characteristic delay in the nodal spike prior to failure (Extended Data Fig. 2a-d). Simulating spike propagation by applying a brief current pulse to mimic the current arriving from an upstream node (and FP) yielded similar results, with full spikes evoked at rest, but hyperpolarization leading to a spike delay and then failure (Extended Data Fig. 3). Large depolarizations inactivated spikes, though outside of the physiological range (> - 50 mV, Extended Data Fig. 3b-c).

Nodal spike facilitation by GABA.

Since sensory axons are tonically depolarized by spontaneous GABA activity¹⁶, we wondered whether this GABA aids propagation. Blocking extrasynaptic α5 GABA_A receptors (with L655708) or all GABA_A receptors (with gabazine) increased the incidence of spike failure (to ~45% and 65%, respectively; Fig. 2f) and sensitivity to hyperpolarization (Extended Data Fig. 2e-h), without altering overall spike properties (Fig. 2g), implying that spike propagation is highly dependent on nodal GABA

receptors. Application of 5-HT to mimic natural brainstem-derived 5-HT also increased failure (Fig. 2f), likely via its indirect inhibition of GABA_A receptor activity³³.

Nodal spike facilitation by GABA_{axo} neuron activation.

To examine whether GABA_{axo} neurons facilitate spike propagation, we expressed light-sensitive channelrhodopsin-2 (ChR2) in GAD2⁺ neurons in adult GAD2⁻CreER/+;R26^{LSL-ChR2-EYFP} mice (termed GAD2//ChR2-EYFP mice, Fig. 3). A brief light pulse (5 - 10 ms) produced a long-lasting depolarization and spiking in these GABA_{axo} neurons (Fig. 3a), followed by a longer lasting GABA_a-mediated depolarization (PAD) of proprioceptive axons at a monosynaptic latency that was blocked by gabazine (Fig. 3a-b). In these mice, spikes in proprioceptive axons failed with a similar incidence as observed in rats (Figs. 3c-h), but the light-evoked PAD prevented this failure (Fig. 3e-g), similar to direct depolarization. In branches with secure non-failing spikes, light had minor effects (Fig. 3d), but blocking GABA_A receptors again increased the incidence of spike failure (Fig. 3h).

In GAD2//ChR2-EYFP or GAD2//ChR2-EYFP//tdTom mice the EYFP and tdTom reporters labelled GABAergic neurons (Fig. 3k; VGAT⁺, GAD2⁺ and VGLUT1⁻) residing near the central canal and throughout much of the dorsal horn (Fig. 3i-o). These neurons densely innervated the dorsal horn with terminal boutons (Fig. 3j,l,n), and less densely innervated both the ventral horn and dorsal columns (Fig. 3l,m,o), allowing GABAergic innervation of sensory axons along their entire length. They made both synaptic (VGAT⁺) and perisynaptic contacts all along proprioceptive sensory axons, both at nodes and sensory axon terminals on motoneurons (Figs. 3k, and 1e), confirming their identity as GABAaxo neurons.

Computer simulations of branch point failure and rescue by GABA.

To establish that spike failure arises at the branch points where GABA can influence them, we generated a computer simulation of a proprioceptive sensory axon arbour in the spinal cord³⁴. With simulated DR stimulation, spike failure occurred distal to complex branch points (at nodes N2 and N3 in Extended Data Fig. 4a-b) that had associated increases in net conductance, which shunted the nodal currents. Simulated nodal GABAA receptor activation rescued these failed spikes, with increasing GABAA activation (g_{GABA}) preventing more branch point failures (Extended Data Fig. 4c). In contrast, when we moved all these GABAA receptors to the terminals, then their activation did not rescue failed spikes (Extended Data 4d). This is because GABAA-induced depolarizations (PAD) were attenuated

sharply with distance ($\lambda s \sim 90 \mu m$); so only nodal, and not terminal, induced PAD was visible at the dorsal columns (Extended Data Fig. 4a,g-h), in agreement with previous terminal recordings¹⁶.

Spike facilitation by sensory evoked GABA_{axo} activity

We next examined whether natural activation of GABA_{axo} neurons affects proprioceptive axon conduction. GABA_{axo} neurons are indirectly activated by sensory activity via two variants of a trisynaptic circuit, where sensory axons drive excitatory neurons that activate GABA_{axo} neurons and cause PAD: one driven by cutaneous afferents and the other by proprioceptive afferents (Extended Data Figs. 5a and 6a)¹⁶. As expected, following DR stimulation these circuits caused fast synaptic and slower extrasynaptic GABA_A receptor mediated depolarizations of proprioceptive axons (termed sensory-evoked phasic PAD and tonic PAD, respectively¹⁶) that were blocked by GABA_A antagonists, and mimicked by optogenetic activation of GABA_{axo} neurons (Fig. 4a-d).

Like with direct GABA_{axo} activation, spike propagation failure was prevented by sensory-evoked phasic PAD, regardless of whether the failure was spontaneous (Figs. 4e-f,h), 5-HT-induced (Fig. 4h), or repetition-induced (Extended Data Fig. 6b-f). The latter is particularly important because sensory axons naturally fire at high rates, where they are vulnerable to spike failure (Fig. 2e-f). This action of phasic PAD was abolished by gabazine but not L655935, supporting a synaptic origin (Fig. 4h). Slow extrasynaptic GABAergic depolarization (tonic PAD; L655935-sensitive¹⁶) further facilitated spike propagation (Fig. 4g), especially as it built up with repeated DR stimulation (at 1 Hz; Extended Data Fig. 5b). Cutaneous (Extended Data Fig. 5), proprioceptive (Extended Data Fig. 6) or mixed afferent (Fig. 4e-h) -evoked PAD all helped prevent spike failure.

In secure non-failing axon branches sensory-evoked PAD (or optogenetic GABA_{axo} activation) sped up the spikes and lowered their threshold (rheobase current; Fig. 3d and Extended Data Fig. 7a-d), as predicted from computer simulations (Extended Data Fig. 4e). Importantly, spike height was only slight reduced during PAD (~1% or 1 mV) indicating that nodal GABA_A receptor conductances have minimal shunting action (Fig. 3 and Extend Data Fig. 7a-d).

Failure of axon conduction to motoneurons and rescue by PAD.

To quantify the overall failure of spikes to conduct from the DR to the sensory axon terminals we measured whether axon branches not conducting during failure were *not* refractory to subsequent stimulation with a microelectrode in the ventral horn (Extended Data Fig. 8). This method indicated that

about 50 - 80% of sensory axons failed to conduct to their ventral terminals under resting conditions, especially in long axons, whereas sensory-evoked PAD decreased failure to < 30%. Similar conclusions were reached by directly recording the extracellular afferent volley in the ventral horn produced by the spikes propagating from a DR stimulation to the motoneurons, which was consistently increased by PAD (Extended Data Fig. 9).

Facilitation of sensory feedback to motoneurons by nodal GABA_A receptors.

To examine the functional role of nodal GABA in regulating sensory feedback to motoneurons, we recorded monosynaptic excitatory postsynaptic potentials (EPSPs) from motoneurons in response to proprioceptive sensory axon stimulation (Fig. 5a). This EPSP was inhibited by optogenetically silencing GABA_{axo} neurons with light in mice expressing archaerhodopsin-3 (Arch3, induced in GAD2^{CreER/+};R26^{LSL-Arch3-GFP} mice; abbreviated GAD2//Arch3, Fig. 5a-b,d), consistent with a tonic GABA_A receptor tone facilitating spike transmission in axons. Likewise, the EPSP was reduced when sensory axon conduction was reduced by blocking endogenous GABA_A receptor tone with antagonists, despite increasing motoneuron and polysynaptic reflex excitability (minimized with APV, Fig. 5c,d). GABA_B antagonists slightly increased the EPSP, suggesting a tonic GABA_B-mediated presynaptic inhibition (Fig. 5d), though much smaller than the tonic GABA_A-mediated nodal facilitation that dominates when all GABA was reduced (in GAD2//Arch3 mice).

Consistent with nodal GABA_A receptors (and PAD) facilitating axon conduction, the monosynaptic EPSP was facilitated during, but not after, depolarizing proprioceptive axons (evoking PAD) with an optogenetic activation of GABA_{axo} neurons in GAD2//ChR2 mice (10 ms *light conditioning stimulation*; Fig. 5e-f). The EPSP was also facilitated by naturally activating GABA_{axo} neurons by a *sensory conditioning stimulation*, including with a conditioning stimulation of cutaneous and/or proprioceptive afferents (Extended Data Fig. 10a,b,e). The latter indicates that proprioceptive activity primes subsequent proprioceptive reflex transmission (*self-facilitation*). GABA_A receptor antagonists (gabazine), but not GABA_B antagonists (CGP55845), blocked the EPSP facilitation with sensory (Extended Data Fig. 10e) or light (Fig. 5f) conditioning.

The facilitation of the EPSP by conditioning-evoked PAD arose from axonal GABAA receptors, rather than from postsynaptic actions on the motoneurons, since it occurred with weak conditioning stimuli that produced only a transient background postsynaptic depolarization that terminated before the EPSP testing (at 60 ms; Figs. 5e, Extended Data Fig. 10b,g), followed by a slight hyperpolarization that if

anything reduced the EPSP (shunting the synaptic current, Extended Data Fig. 10h). Increasing the DR conditioning intensity produced large background depolarizing conductances in the motoneurons during the EPSP testing, which led to postsynaptic inhibition of the EPSP (shunting inhibition; Extended Data Fig. 10d,g), masking the effect of nodal facilitation. Importantly, sometimes PAD itself induced afferent spikes (Extended Data Fig. 7e; termed DRR spikes), and following these spikes, the EPSP was always smaller than when these spikes were not present (n = 8/8 mice, not shown). This is because these DRR spikes themselves triggered EPSPs, leading to a post activation depression, as noted by Eccles⁹, and thus we minimized DRR activity by keeping the conditioning-evoked PAD small.

Sensory conditioning was particularly effective when it was repeated to mimic natural firing, which increased tonic PAD for minutes (Fig. 5g). This facilitated the EPSP for ~3 min after a brief fast DR repetition (200 Hz, 0.5 s conditioning, Fig. 5i, Extended Data Fig. 10e, *Tonic*), and ~1 min after slower repetition (0.1 Hz, 2 min conditioning, Extended Data Fig. 10e, *After effect*), both long outlasting postsynaptic effects from each conditioning pulse (< 1 s). This was blocked by L655708 or gabazine (Extended Data Fig. 10e).

Increases in the probability of unitary EPSPs by nodal facilitation.

We often noticed large all-or-nothing EPSPs (unitary EPSPs) spontaneously fluctuating on and off during repeated EPSP testing, leading to discrete large changes in the total EPSP size and time course (Fig. 5j-k). We thought this might be due to spontaneous branch point failures, rather than quantal changes in transmitter release that produce much smaller fluctuations³⁵, as previously suggested²⁸. Indeed, when we increased the axon conduction by activating the GABA_{axo} neurons and PAD (via a cutaneous conditioning train) the *probability* of unitary EPSPs occurring increased (Fig. 5k-l), and this sometimes recruited further large unitary EPSPs (Fig. 5k). In contrast, the *size* of the underlying unitary EPSP was not increased by this conditioning (Fig. 5j-l), ruling out decreases in terminal presynaptic inhibition or postsynaptic inhibition contributing to the increased overall EPSP (Fig. 5i,l).

Facilitation of sensory feedback to motoneurons by GABA_{axo} neurons in awake mice.

To determine whether GABA_{axo} neurons increase sensory feedback to motoneurons in *awake* mice we activated these neurons with light applied through a window chronically implanted over the spinal cord of GAD2//ChR2 mice (Fig. 6a), and assessed the monosynaptic reflex (MSR) recorded in tail muscles in response to nerve stimulation (counterpart of EPSPs). As expected, the MSR was facilitated by a conditioning light pulse, but only during, and not after, the expected time of phasic PAD induced on

sensory axons (Fig. 6b-d,j). This light-induced *facilitation* occurred both at rest and when there was a background voluntary contraction, with the latter matched with and without light, again ruling out postsynaptic depolarization related differences in MSR (Fig. 6d). Light alone caused a brief pause in ongoing EMG (~30 ms post-light; Fig. 6b), indicative of postsynaptic inhibition, which masked nodal facilitation at short intervals.

Facilitation of sensory feedback to motoneurons during PAD in humans and rats

Finally, to estimate the role of GABA_{axo} neurons in humans (and awake rats) we employed the sensory-evoked depolarization of proprioceptive axons by GABA_{axo} neurons (sensory-evoked PAD; Fig. 4), which is known to occur in humans³⁶. For this we recorded the MSR in humans in the soleus muscle in response to tibial nerve stimulation (Fig. 7a). Increasing GABA_{axo} neuron activity with a brief cutaneous stimulation increased the MSR (Fig 7bi, d and Fig. 6e-g) during a period consistent with nodal facilitation by PAD (30 – 200 ms post stimulation; in humans and rats, Fig. 7biii and Fig. 6j). We again kept the conditioning stimulation small enough to not change the background EMG or single motor unit (MU) firing (Fig. 7biii, Fig 6f-g) to rule out postsynaptic actions. When we instead increased PAD by a proprioceptive conditioning in humans (via muscle TA vibration) the soleus MSR was inhibited (for up to 200 ms; Fig. 7ci-ii), as previously reported³⁷. However, the vibration alone inhibited the ongoing MU discharge (Fig. 7ciii), implying that this MSR inhibition was caused in part by postsynaptic inhibition, rather than PAD-mediated presynaptic inhibition³⁷.

Blocking GABA_A receptor tone (in rats) decreased the MSR, at matched levels of background EMG (Fig. 6h), suggesting a spontaneous tonic PAD facilitating the MSR. In rats and humans repeated cutaneous conditioning stimulation (trains) to induce a buildup in this tonic PAD caused an associated buildup of the MSR that outlasted the conditioning and its postsynaptic actions by many seconds (after effect; Fig. 7d,e; Fig. 6i).

Importantly, the probability of a single MU contributing to the human MSR was increased by cutaneous conditioning (Fig. 7fi-ii). This occurred without an increase in the estimated EPSP amplitude or rise time (PSF; see Methods; Fig. 7Fiii) or change in the MU firing prior to the MSR testing (Fig. 7Fiv; motoneuron not depolarizing closer to threshold), consistent with an increased probability of unitary ESPSs, as in rats (Fig. 5).

Discussion

Following the pioneering studies of Eccles on inhibition of the monosynaptic connection from sensory axons to motoneurons⁹, the concept of presynaptic inhibition of axon terminals has stood as a cornerstone of our understanding of mammalian brain and spinal cord function¹⁰. While presynaptic inhibition has never been directly confirmed in these sensory axons, recordings from invertebrate sensory axons have firmly established the idea that terminal GABAA receptor-mediated depolarizations (PAD) can cause conductance increases (shunts) or sodium channel inactivation that inhibit transmitter release^{22,24}. Thus, our finding that GABA_A receptors are located too far from the axon terminals to influence terminal depolarizations (relative to short λ_s) or cause presynaptic inhibition of transmitter release onto motoneurons (EPSPs not inhibited) had not been anticipated. Direct recordings from terminal boutons of other axon types in the mammalian brain (e.g. Calyx of Held) have shown that terminal GABAA or glycine receptors sometimes cause presynaptic inhibition, and at other times facilitate transmitter release, depending on their action on terminal calcium and potassium channels^{21,22,38}. However, these terminal actions of GABA are very different from the actions of GABA on the nodes that we have uncovered, where GABAA receptors near sodium channels (nodes) and associated branch points help prevent conduction failure by bringing nodes closer to spike threshold. We found that without nodal GABA provided by GABA_{axo} neurons, spike transmission fails in many central branches of large myelinated sensory axons, leaving large silent branches, depending on the branching structure and prior history of activity (frequency). Thus, neuronal circuits that control GABA_{axo} neurons serve to aid branch conduction via nodal facilitation. This concept of nodal facilitation may generalize to other large central axons (e.g. pyramidal cells) that are innervated by GABAergic neurons, branch extensively (and so may fail), and have depolarizing actions of GABA^{20,22,39,40}, allowing selective recruitment of specific axon branches and functional pathways, especially for high frequency firing.

Sensory driven GABA_{axo} circuits (and PAD) are experimentally convenient, since they allow us to estimate how sensory transmission to motoneurons is modulated with GABA_{axo} neuron activity in humans. Specifically, the expected PAD evoked by cutaneous conditioning is associated with a potent reflex facilitation in humans and awake rodents. This suggests that a substantial ongoing spike failure (prior to facilitation) that can be alleviated by GABA_{axo} activity (PAD). Indeed, we found that during PAD the *probability* of EPSPs occurring (and MU firing) is increased without changing the EPSP amplitude (estimated by PSFs in humans). The latter rules out changes in presynaptic inhibition with

PAD that *grades* the EPSP size, including ruling out previous arguments that MSR facilitation by cutaneous conditioning is due to a removal of presynaptic inhibition^{41,42}.

A pressing question that remains is how can nearly a century of research on sensory transmission and presynaptic inhibition be reconciled with GABA-mediated nodal facilitation and reflex facilitation (Table S1)? Sensory axon conduction failure has repeatedly been noted from indirect observations²⁸⁻³⁰, but GABAA receptors and PAD were previously thought to cause, rather than prevent, conduction failure³⁰, even though computer simulations showed physiological GABA levels unlikely to block spike propagation³⁴, as we confirmed. Furthermore, the fundamental assumption that GABA_A receptors cause presynaptic inhibition that reduces transmitter release from sensory axons was from the outset circumspect, based mainly on the observation that a conditioning stimulation (on a flexor nerve) caused an inhibition of the MSR (evoked in extensor muscles) that was somewhat correlated to the time-course of PAD caused by this conditioning (in extensor afferents)⁹. However, in retrospect this PAD is too brief to account for the much longer (up to 1 s) inhibition caused by this conditioning^{9,11,43}, and GABAB receptor antagonists block much of this MSR inhibition^{17,43}. This fits with GABA_B receptors being at the terminals (Fig. 1) and primarily responsible for presynaptic inhibition in proprioceptive axons, as in other neurons^{21,22}, though further study of GABA_B receptor function is now needed. This predominant GABA_B action in proprioceptive axon terminals does not rule out GABA_A-mediated presynaptic inhibition in other sensory axons that have dense terminal GABAA receptor expression, such as in cutaneous afferents¹⁶.

Anatomical studies have shown that GABA_{axo} neuron activation is inevitably accompanied by some postsynaptic inhibition, since most GABA_{axo} contacts on afferent terminals also contact motoneurons, in a triad^{8,44}. Indeed, we find that GABA_{axo} neuron activation produces an inhibition of motoneurons (Fig. 6b) and associated MU firing (Fig. 7c) that masks, and at times overwhelms, the facilitation of the MSR by nodal GABA_A receptors (as with muscle vibration). This postsynaptic inhibition was likely previously mistaken for presynaptic inhibition, though this needs to be re-examined³⁷. The argument that presynaptic inhibition with conditioning should be evident from reductions in the EPSP without changing its time course⁴⁵ now seems untenable, especially as unitary EPSPs differ markedly in shape and conditioning increases the number of unitary EPSPs contributing to the EPSP, as different axon branches are recruited (Fig. 5k)²⁸.

Early on Barron and Matthews²³ and later others^{7,16} established that sensory-evoked PAD (or light-evoked) excites axons by directly inducing spiking, including spikes in the sensory axons mediating the MSR itself, raising a further contradiction with presynaptic inhibition. While these PAD-triggered spikes only sometimes fully propagate antidromically out the DR⁴⁶, they are more likely to conduct orthodromically¹⁶ where they activate the motoneurons^{7,9,47}, making these axons and their motoneuron synapse refractory to subsequent testing⁹. This contributes to a long-lasting *post-activation depression* of the MSR pathway that is GABA_A-mediated (sensitive to GABA_A antagonists, like PAD) and is thus readily mistaken for GABA_A-mediated presynaptic inhibition^{7,17,18,37}.

Functionally, nodal facilitation and regulation of branch point failure by GABA_{axo}-driven GABA_A receptors acts like a global switching system that recruits entire silent sensory or motor circuits. This works in concert to terminal presynaptic inhibition (including GABA_B receptor action) that locally fine tunes reflex gains, to optimize the stability and compliance of movement^{3,4,7}. The direct activation of GABA_{axo} neurons (and PAD) by cortical (CST) and spinal (CPG) circuits^{12,13,48}, and inhibition by the brainstem (5-HT)^{33,49}, suggests that nodal facilitation is under explicit central control during reaching and locomotion (Fig. 2). The widespread action of PAD (occurring simultaneously over many spinal segments)^{11,16} implies that nodal facilitation acts over large regions of the spinal cord to ready sensory axons for action during cortical, spinal or sensory evoked activity, reminiscent of the Jendrassik maneuver⁵⁰, ensuring that adequate sensory feedback aids postural stability and walking. More generally, our results imply that each axonal branch point has the capacity to function separately (like separate neurons), depending on their GABAergic innervation, increasing the complexity of sensory processing in the spinal cord.

Methods

Adult mice, rats and humans used.

Recordings were made from large proprioceptive group Ia sensory afferents, GABAergic neurons, motoneurons and muscles in adult mice (2.5 – 6 months old, both female and male equally; strains detailed below) and rats (3 - 8 months old, female only, Sprague-Dawley). All experimental procedures were approved by the University of Alberta Animal Care and Use Committee, Health Sciences division. Recordings were also made from the soleus muscle of neurologically intact adult humans (female and male equally), aged 21 to 58, with written informed consent prior to participation. Experiments were approved by the Health Research Ethics Board of the University of Alberta (Protocols 00023530 and 00076790) and conformed to the Declaration of Helsinki. No effects of sex were noted and data from both sexes were combined for analysis.

Mice used for optogenetics and imaging.

We evaluated GABAergic neurons in a strain of mice with Cre expressed under the endogenous *Gad2* promotor region. *Gad2* encodes the Glutamate decarboxylase 2 enzyme GAD2 (also called GAD65), which is unique to axoaxonic contacting GABAergic neurons that project to the ventral horn, whereas all GABAergic neurons express GAD1⁶. These GAD2⁺ neurons were activated or inhibit optogenetically using channelrhodopsin-2 (ChR2)^{51,52} or archaerhodopsin-3 (Ach3)^{53,54}, respectively. The following mouse strains were employed (Supplementary Table 2):

- 1) $Gad2^{tm1(cre/ERT2)Zjh}$ mice (abbreviated $Gad2^{CreER}$ mice; The Jackson Laboratory, Stock # 010702; CreER^{T2} fusion protein expressed under control of the endogenous Gad2 promotor)⁵⁵,
- 2) B6;129S-*Gt(ROSA)*26Sor^{tm32(CAG-COP4*H134R/EYFP)Hze} mice (abbreviated R26^{LSL-ChR2-EYFP} mice; The Jackson Laboratory, Stock # 012569; ChR2-EYFP fusion protein expressed under the R26::CAG promotor in cells that co-express Cre because a loxP-flanked STOP cassette, LSL, prevents transcription of the downstream ChR2-EYFP gene)⁵⁶,
- 3) B6.Cg-*Gt*(*ROSA*)26Sor^{tm14(CAG-tdTomato)Hze and B6.Cg-*Gt*(*ROSA*)26Sor^{tm9(CAG-tdTomato)Hze mice (abbreviated R26^{LSL-tdTom} mice; The Jackson Laboratory, Stock # 007914 and #007909; tdTomato fluorescent protein expressed under the R26::CAG promotor in cells that co-express Cre)⁵⁷,}}
- 4) B6;129S-*Gt*(*ROSA*)26Sor^{tm35.1(CAG-aop3/GFP)Hze} mice (abbreviated R26^{LSL-Arch3-GFP} mice; The Jackson Laboratory Stock # 012735; Arch3-GFP fusion protein expressed under the R26::CAG promotor in cells that co-express Cre)⁵⁶, and

5) B6;129S-*Slc17a7*^{tm1.1(cre)Hze} mice (abbreviated VGLUT1^{Cre} mice; The Jackson Laboratory, Stock # 023527; Cre protein expressed under control of the endogenous *Vglut1* promotor; kindly donated by Dr. Francisco J. Alvarez)⁵⁸.

Heterozygous GAD2^{CreER} mice (i.e., GAD2^{CreER/+} mice) were crossed with homozygous reporter strains to generate GAD2^{CreER/+}; R26^{LSL-ChR2-EYFP}, GAD2^{CreER/+}; R26^{LSL-tdTom} and GAD2^{CreER/+}; R26^{LSL-Arch3-GFP} mice that we abbreviate: GAD2//ChR2, GAD2//tdTom and GAD2//Arch3 mice. Offspring without the GAD2^{CreER} mutation, but with the effectors ChR2, Arch3 or tdTom were used as controls. We also used mice bred by crossing homozygous VGLUT1^{Cre} mice with R26^{lsl-tdTom} reporter mice to obtain mice with VGLUT1 labelled sensory axons ⁵⁹.

CreER is an inducible form of Cre that requires tamoxifen to activate ⁶⁰, which we applied in adult mice to prevent developmental issues of earlier induction of Cre. Specifically, mice were injected at 4 - 6 weeks old with two doses of tamoxifen separated by two days, and studied > 1 month later, long after washout of tamoxifen. Each injection was 0.2 mg/g wt (i.p.) of tamoxifen dissolved in a corn oil delivery vehicle (Sigma C8267). These tamoxifen-treated mice were denoted GAD2//ChR2+ and GAD2//Arch3+, and non treated mice were used as controls and denoted GAD2//ChR2- and GAD2//Arch2-. For all mice, genotyping was performed according to the Jackson Laboratories protocols by PCR of ear biopsies using primers specific for the appropriate mutant and wild type alleles for each of the mouse lines (see Key Resources Table for primer details).

Ex vivo recording from axons and motoneurons in whole adult spinal cords.

Mice or rats were anaesthetized with urethane (for mice 0.11 g/100 g, with a maximum dose of 0.065 g; and for rats 0.18 g/100 g, with a maximum dose of 0.45 g), a laminectomy was performed, and then the entire sacrocaudal spinal cord was rapidly removed and immersed in oxygenated modified artificial cerebrospinal fluid (mACSF), as detailed previously ⁶¹⁻⁶³. This preparation is particularly useful as the small sacrocaudal spinal cord is the only portion of the adult spinal cord that survives whole ex vivo, allowing axon conduction to be assessed along large distances. Further, this segment of cord innervates the axial muscles of the tail that are readily assessable for reflex recording in awake animals, and has proven to be a useful model of motor function in normal and injured spinal cords ^{62,64}. Spinal roots were removed, except the sacral S3, S4 and caudal Ca1 ventral and dorsal roots on both sides of the cord. After 1.5 hours in the dissection chamber (at 20° C), the cord was transferred to a recording chamber containing normal ACSF (nACSF) maintained at 23 - 32°C, with a flow rate > 3 ml/min. A one-hour

period in nACSF was given to wash out the residual anaesthetic prior to recording, at which time the nACSF was recycled in a closed system. The cord was secured onto tissue paper at the bottom of a rubber (Silguard) chamber by insect pins in connective tissue and cut root fragments. The dorsal surface of the cord was usually oriented upwards when making intracellular recording from afferents in the dorsal horn, whereas the cord was oriented with its left side upwards when making recordings from motoneurons or afferent terminals in the ventral horn. The laser beam used for optogenetics was focused vertically downward on the GAD2 neurons, as detailed below.

Optogenetic regulation of GABA_{axo} neurons.

The GAD2//ChR2 or GAD2//Arch3 mice were used to optogenetically excite or inhibit GAD2+ neurons (with 447 nm D442001FX and 532 nM LRS-0532-GFM-00200-01 lasers from Laserglow Technologies, Toronto), respectively, using methods we previously described 65 . Light was derived from the laser passed through a fibre optic cable (MFP_200/220/900-0.22_2m_FC-ZF1.25 and MFP_200/240/3000-0.22_2m_FC-FC, Doric Lenses, Quebec City) and then a half cylindrical prism the length of about two spinal segments (8 mm; 3.9 mm focal length, Thor Labs, Newton, USA,), which collimated the light into a narrow long beam (200 μ m wide and 8 mm long). This narrow beam was focused longitudinally on the left side of the spinal cord roughly at the level of the dorsal horn, to target the epicentre of GABAaxo neurons, which are dorsally located (Fig. 3). ChR2 rapidly depolarizes neurons 52 , and thus we used 5-10 ms light pulses to activate GABAaxo neurons, as confirmed by direct recordings from these neuron (see below). Light was always kept at a minimal intensity, 1.1x T, where T is the threshold to evoke a light response in sensory axons, which made local heating from light unlikely. Arch3 is a proton pump that is activated by green light, leading to a hyperpolarization and slowly increased pH (over seconds), both of which inhibit the neurons 52,66 . Thus, we used longer light pulses (\sim 200 ms) to inhibit GABAaxo neurons.

To directly confirm the presence of functional GAD2 expression in neurons (GABA_{axo} neurons) we recorded from them with similar methods and intracellular electrodes used to record from motoneurons (see below). Electrodes were advanced into these cells through the dorsal horn (with the dorsal surface oriented upwards), and their identity established by a direct response to light activation of the ChR2 construct (5 – 10 ms light pulse, 447 nm, mW; in GAD2//ChR2 mice), without a synaptic delay (<1 ms) and continued light response after blocking synaptic transmission.

Dorsal and ventral root stimulation.

Dorsal and ventral roots (DR and VR) were mounted on silver-silver chloride wires above the nASCF of the recording chamber and covered with grease (a 3:1 mixture of petroleum jelly and mineral oil) for monopolar stimulation 16,64,67 . This grease was surrounded by a more viscous synthetic high vacuum grease to prevent oil leaking into the bath flow. Bipolar stimulation was also used at times to reduce the stimulus artifact during recording from ventral roots (detailed below). Roots were stimulated with a constant current stimulator (Isoflex, Israel) with short pulses (0.1 ms). Note that proprioceptive afferents are selectively activated by low intensity DR stimulation (1.1 – 1.5 x threshold, T) and cutaneous afferents are additionally activated by higher intensity DR stimulation (2 – 3xT). DRs were dissected to be as long as possible, and the distal end of this root was stimulated, so it was ~20 mm way from the spinal cord. In this way the DR stimulation site itself (at wire, and threshold for stimulation) could not be affected by axonal depolarizations in the spinal cord, since dorsal root potentials from spinal events (PAD) are only observed very close to the cord (within a few mm, see below), and drop exponentially in size with distance 16 .

Intracellular recording from sensory axon branches in the dorsal horn.

Electrode preparation and amplifier. Recording from fine afferent collaterals in the spinal cord without damaging them or disturbing their intracellular milieu required specialized ultra-sharp intracellular electrodes modified from those we developed for motoneuron recording ⁶¹. That is, glass capillary tubes (1.5 mm and 0.86 mm outer and inner diameters, respectively; with filament; 603000 A-M Systems; Sequim, USA) were pulled with a Sutter P-87 puller (Flaming-Brown; Sutter Instrument, Novato, USA) set to make bee-stinger shaped electrodes with a short relatively wide final shaft (~1 mm) that tapered slowly from 30 to 3 µm over its length, and then abruptly tapered to a final tip over the final 20 µm length. The tip was subsequently bevelled to a 100 nm hypodermic-shaped point, as verified with electron microscope images (Harvey et al. 2006). This very small tip and wide shaft gave a combination of ease of penetrating axons in dense adult connective tissue, and good current-passing capabilities to both control the potential and fill the axons with neurobiotin. Prior to beveling, electrodes were filled through their tips with 2 M K-acetate mixed with varying proportions of 2 M KCl (to make intracellular Cl- concentrations ranging of 0, 100, 500, and 1000 mM) or 500 mM KCl in 0.1 Trizma buffer with 5 -10% neurobiotin (Vector Labs, Birmingame, USA). They were then beyeled from an initial resistance of 40 - 150 M Ω to 30 - 40 M Ω using a rotary beveller (Sutter BV-10). GABAergic chloride-mediated potentials (PAD) were the same with different concentrations of KCl, without passing large amounts of negative current, as we have previously detailed ¹⁶, indicating that the ultra-sharp tips impeded passive fluid exchange between the electrode and intracellular milieu, with in particular electrode Cl⁻ not

affecting the axon; thus, recordings were mostly made with electrodes with 1 M K-acetate and 1 M KCl, when not filling cells with neurobiotin.

Intracellular recording and current injection were performed with an Axoclamp2B amplifier (Axon Inst. and Molecular Devices, San Jose, USA). Recordings were low pass filtered at 10 kHz and sampled at 30 kHz (Clampex and Clampfit; Molecular Devices, San Jose, USA). Sometimes recordings were made in discontinuous-single-electrode voltage-clamp (gain 0.8 –2.5nA/mV; for Ca PICs) or discontinuous-current-clamp modes (switching rate 7 kHz), as indicated (the latter only when injecting current, for example during recording of input resistance or the voltage dependence of spikes).

Axon penetration. Electrodes were advanced into myelinated afferents with a stepper motor (Model 2662, Kopf, USA, 10 µm steps at maximal speed, 4 mm/s), usually at the boundary between the dorsal columns and dorsal horn gray matter. Extracellular tissue (especially myelin in the white matter) often impeded and blocked the electrode tip following a forward step, as determined by an increase in resistance to small current pulses passed from the tip of the electrode (20 ms, -0.3 nA, 1 Hz), and this was cleared with a brief high frequency current (from capacitance overcompensation buzz) and moving backwards slowly, the latter which helped prevent tissue dimpling. Prior to penetrating afferents, we recorded the extracellular (EC) afferent volley following dorsal root (DR) stimulation (0.1 ms pulses, 3xT, threshold, where T = -3 uA, repeated at 1 Hz), to determine the minimum latency and threshold of afferents entering the spinal cord. The group Ia afferent volley occurs first with a latency of 0.5 - 1.0 ms, depending on the root length (which were kept as long as possible, 10 - 20 mm), corresponding to a conduction velocity of about 16 - 24 m/s, as previously described for in vitro conduction at 23 C ^{16,68}. When a forward step penetrated an axon, small slow movements were made to stabilize the recordings. Penetrations were usually in the myelinated portion of the axon between nodes, rather than at nodes, because the chance of penetrating a node is low since they only make up a small fraction of the total axon length (Fig. 1). The spikes from the two nodes adjacent to the electrode were readily detected separately when testing for the spike threshold with current injection pulses (20 ms; rheobase test), because just at threshold the current sometimes evoked a spike from just one node and not the other, which usually halved the total spike height, consistent with the penetration being about halfway between the two nodes.

Proprioceptive afferent identification. Upon penetration, afferents were identified with direct orthodromic spikes evoked from DR stimulation. We focused on the lowest threshold (T)

proprioceptive group Ia afferents, identified by their direct response to DR stimulation, very low threshold (< 1.5 x T), short latency (group Ia latency, coincident with onset of afferent volley), and antidromic response to ventral horn afferent terminal microstimulation (~ 10 μ A stimulation via tungsten microelectrode to activate Ia afferent terminals; tested in some afferents, detailed below)¹⁶. Clean axon penetrations without injury occurred abruptly with a sharp pop detected on speakers attached to the recorded signal, the membrane potential settling rapidly to near – 70 mV, and > 70 mV spikes readily evoked by DR stimulation or brief current injection pulses (1 – 3 nA, 20 ms, 1 Hz). Sensory axons also had a characteristic >100 ms long depolarization following stimulation of a dorsal root (primary afferent depolarization, PAD, at 4 - 5 ms latency, detailed below) and short spike afterhyperpolarization (AHP ~ 10 ms), which further distinguished them from other axons or neurons. Injured axons had higher resting potentials (> - 60 mV), poor spikes (< 60 mV) and low resistance (to current pulse; $R_m < 10 \text{ M}\Omega$) and were discarded.

Quantification of spike conduction failure in the dorsal horn: failure potentials (FPs). Sometimes healthy intracellular penetrations were made into a sensory axon branch (e.g. < -60 mV rest, large PAD), but dorsal root stimulation did not evoke a full spike, even though a full > 60 mV spike could be readily evoked by intracellular current injection. Instead, DR stimulation evoked a partial spike at the latency and threshold of group Ia afferents, indicating that this was a branch of a Ia afferent that failed to fully conduct spikes to the electrode, with only the passively attenuated spike from the last node to spike prior to conduction failure recorded at the electrode (failure potential, FP; also referred to as electronic residue by Luscher⁶⁹). The size of the FP reflected how far away the spike failure occurred, with spatial attenuation corresponding to a space constant of about 90 µm (see Results), and so FPs became exponentially smaller with distance from failure and undetectable when many mm away (nodes separated by about 50 µm). Occasionally axons were penetrated with undetectable DR evoked spikes or FPs, but otherwise they had characteristics of a Ia afferent (PAD, R_m similar). These were likely afferents with FPs too distal to detect, but were usually excluded from the main analysis to avoid ambiguity, though this underestimates the incidence of failure. However, some of these axons exhibited short latency, low threshold DR spikes when depolarized by a prior DR stimulation (PAD) of an adjacent DR, in which case they were unequivocally Ia afferents and included in the analysis (Fig. 4f).

Both during extracellular and intracellular recording the group Ia afferent volley (small negative field) was observed as the first event after DR stimulation (the latter subthreshold to a spike), though this was usually small in relation to intracellular events and ignored. However, this was sometimes removed

from the intracellular record by subtracting the extracellular potential recorded just outside the same axon to determine the actual transmembrane potential ¹⁶. This was necessary to see the very smallest FPs following DR stimulation in some afferents, as the negative volley from other nearby afferents obscured the FPs.

After quantifying the axons spikes and conduction failures (FPs) under resting conditions, we then examined the changes in spike conduction with changes in membrane potential induced by either directly injecting current into axons or inducing GABA-mediated changes in membrane potential by pharmacological methods, optogenetic methods (activating ChR2 on GABA_{axo} neurons to induce PAD) or more naturally evoking PAD with a DR stimulation.

Neurobiotin filling of axons. Some of the proprioceptive afferents that we recorded intracellularly were subsequently filled with neurobiotin by passing a very large positive 2 - 4 nA current with 90% duty cycle (900 ms on, 100 ms off) for 10 - 20 min. The identity of group Ia proprioceptive afferents were then confirmed anatomically by their unique extensive innervation of motoneurons ¹⁶. Prior to penetrating and filling axons with neurobiotin filled electrodes, a small negative holding current was maintained on the electrodes to avoid spilling neurobiotin outside axons.

Quantification of spike conduction failure in the ventral horn

Wall's method. To measure whether spikes fail during propagation to their fine terminals in the ventral horn we examined whether failed axon segments were relatively less refractory to activation after spike conduction failure, using a double pulse method adapted from Wall 30,70 . The essence of the method is that after DR activation all nodes that generate spikes become relatively refractory for a few ms, whereas nodes that fail to spike are not refractory to activation. Thus, a microelectrode placed near these failing nodes more readily activates them if they fail rather than generate spikes with DR stimulation and orthodromic conduction. For this we placed a tungston microelectrode (12 M Ω , #575400, A-M Systems, Sequim, USA) in the ventral horn near the axons terminals on motoneurons, to activate the branches/nodes of the axon projecting to the motoneuron that may have failed (VH stimulation).

Spikes from VH or DR stimulation were recorded intracellularly in a proprioceptive axon penetrated in the dorsal columns directly above the VH stimulation site or in an adjacent segment, with two combinations of double axon stimulations. First, we applied two rapidly repeated VH stimuli (VH

doublet; two 0.1 ms pulses) at a \sim 4 ms interval to make the axon relatively refractory to stimulation and determine both the threshold current to activate the first spike (T_{VHI} , with VH1 stimulation) and the higher threshold current to overcome this the inactivation and generate a second spike (T_{VH2} , with VH2 stimulation). Second, we repeated this double spike activation, but with the first activation from a suprathreshold DR stimulation (at 1.5x DR threshold) and the second from a VH stimulation at the T_{VH2} intensity from B (DR-VH pair). In this case the VH stimulation readily activates the axon spike if the orthodromic DR evoked spike does not propagate to the ventral horn, leaving the silent portion of the axon non refractory. Accordingly, we also determined the threshold current to activate the VH after the DH in this arrangement (termed $T_{DR,VH}$), which was lower than T_{VH2} . For comparison to the spike inactivation with VH doublets, we adjusted the DR-VH pair timing slightly so that the pairs of spikes (or expected spikes, at vertical lines) are separated by the same interval (\sim 4 ms) when they reach the recording site, to compensate for DR conduction delays. The putative spike failure with DR stimulation happens at a node somewhere between the recording site and the VH, because we only studied axons that securely conducted single DR pulses to the recording site, and thus failure was not directly visible.

We quantified the spike failure based on the following considerations: If the DR-evoked spike entirely fails to propagate to the VH, then the threshold for subsequently activating the ventral horn $(T_{DR,VH})$ should be the same as the threshold without any prior activation ($T_{VHI} = T_{DR,VH}$), whereas if it does not fail, then the threshold for activating the ventral horn should be the same as with a VH doublet ($T_{VH2} =$ $T_{DR,VH}$). In between these two extreme scenarios, the DR evoked spike may only partially fail to propagate spikes to the ventral horn (by only some of its branches failing or conducting only partially to the VH); in this case $T_{DR,VH}$ should be between T_{VHI} and T_{VH2} , with the difference T_{VH2} - T_{VHI} representing the range of possible thresholds between full failure and full conduction. Thus, overall the failure was quantified as: Conduction failure = $(T_{VH2} - T_{DR,VH})/(T_{VH2} - T_{VH1}) \times 100\%$, which is 100% at full failure and 0% with no failure. This estimate is predicated on the assumption that the failed spikes are only relatively refractory to conduction and increased stimulation can overcome this failure, which is reasonable for the interspike intervals we used, and means that the computed % failure reflects the number of nodes that failed to spike, with more dorsal branch point failures giving more failed nodes. On the other hand, we used interspike intervals that were short enough for the DR stimulation not to evoke PAD that affected the subsequent spike threshold (~4 ms), in contrast to the longer intervals where PAD can help DR doublet firing (DR-DR in Extended Data Fig. 6, ~ 5 - 10 ms).

Extracellular recording from sensory axon terminals. To directly record spike conduction in proprioceptive afferent terminal branches in the VH we used our intracellular glass pipette electrode (\sim 30 M Ω) positioned just outside these axons (extracellular, EC), to avoid penetration injury in these fine axon branches. The DR was stimulated near threshold (1.1xT) to evoke the EC response in a few single axons near the electrode, and many trials were averaged to remove noise from these small signals (20 – 50 trials at 3 s intervals). The EC field was multiphasic as previously described for other axons ⁷¹⁻⁷³, with a small initial positive field resulting from passively conducted axial current from sodium spikes at distant nodes (closer to the DR; outward current at electrode), some of which fail to propagate spikes to the VH recording site, making this field a measure of conduction failure ^{71,73}. Following this, a larger negative field arises, resulting from spikes arising at nodes near the electrode (inward current), making this negative field a measure of secure conduction. A relatively large stimulus artifact is present prior to these fields, due to the small size of the EC fields themselves, and we truncated this.

We conducted three control experiments to confirm the relation of these EC fields to spike conduction. First, in the dorsal horn where we can readily intracellularly record from large proprioceptive axon branches, we compared intracellular (IC) recordings from axons to EC recordings just outside the same axon, to confirm that the DR evoked spike (IC) arrives at about the time of the negative EC field. Second, we locally applied TTX to the DR near the recording site (10 µl bolus of 100 µM TTX over DR) which eliminated the negative field and left only the initial positive field, confirming that the positive field is from distal nodes upstream of the TTX block, and generated by passive axial current conduction. This is important, since some investigators have argued on theoretical grounds that the positive field can instead result from the closed end electrical properties of axons at their terminals ⁷⁴, rather than spike failure, though others have refuted this 73. Finally, we improved nodal spike conduction by reducing the divalent cations Mg⁺⁺ and Ca⁺⁺ in the bath medium, since divalent cations normally cause a gating or guarding action on the sodium channel, the latter by one charge binding to the membrane and the other raising the local extracellular positive charge, and overall raising the local voltage drop across the channel and its spike threshold⁷⁵. This decreased the failure-related initial positive field and increased the main EC negative field, indicating improved conduction, and again confirming the use of these fields as measures of conduction, similar to previous conclusions for the motor endplate ⁷¹ and mathematical consideration of axon cable properties ⁷⁶.

To quantify the EC fields we estimated the overall conduction to the recording site as:

Conduction Index = nf/(nf + pf) x 100%, where pf and nf are the positive and negative EC field amplitudes. This conduction index approaches 100% for full conduction ($pf \sim 0$) and 0% for no conduction ($pf \sim 0$). The absolute EC field potential amplitudes are highly variable between different recordings sites, and thus are difficult to quantify across animals and sites, whereas this ratio of field amplitudes ($pf \sim 0$) eliminates the variability, and can effectively be viewed as a normalization of the negative field ($pf \sim 0$) by the total field peak-to-peak size ($pf \sim 0$).

Intracellular recording from motoneurons.

The same intracellular glass electrode, stepper motor and amplifier used for recording sensory axons were used for intracellular recording from motoneurons, except that the electrodes were bevelled to a lower resistance (30 M Ω). The electrode was advanced into motoneurons with fast 2 μ m steps and brief high frequency currents (capacitance overcompensation) guided by audio feedback from a speaker. After penetration, motoneuron identification was made with antidromic ventral root stimulation, and noting ventral horn location, input resistance and time constant (> 6 ms for motoneurons) 62 . The monosynaptic excitatory postsynaptic potentials (EPSPs) and associated currents (EPSCs) were measured in motoneurons following stimulation of dorsal roots (at 1.1- 1.5 xT, 0.1 ms, 3 – 10 s trial intervals). These were identified as monosynaptic by their rapid onset (first component), lack of variability in latency (< 1 ms jitter), persistence at high rates (10 Hz) and appearance in isolation at the threshold for DR stimulation (< 1.1xT), unlike polysynaptic EPSPs which varying in latency, disappear at high rates, and mostly need stronger DR stimulation to activate.

Dorsal and ventral root grease gap recording.

In addition to recording directly from single proprioceptive axons and motoneurons, we employed a grease gap method to record the composite intracellular response of many sensory axons or motoneurons by recording from dorsal and ventral roots, respectively, as previously detailed for similar sucrose and grease gap methods, In this case, a high impedance seal on the axon reduces extracellular currents, allowing the recording to reflect intracellular potentials ^{16,76-78}. We mounted the freshly cut roots onto silver-silver chloride wires just above the bath, and covered them in grease over about a 2 mm length, as detailed above for monopolar recordings. Return and ground wires were in the bath and likewise made of silver-silver chloride. Specifically for sensory axons, we recorded from the central ends of dorsal roots cut within about 2 - 4 mm of their entry into the spinal cord, to give the compound potential from all afferents in the root (dorsal roots potential, DRP), which has previously been shown to correspond to PAD, though it is attenuated compared to the intracellular recordings of PAD¹⁶. The

signal attenuation has two reasons. First the voltage PAD is attenuated along the length of nerve in the bath, as detailed in the next paragraph. Second, the grease does not completely remove the extracellular fluid around the nerve, even though we deliberately allowed the nerve to dry for a few seconds before greasing, and this causes a conductance that shunts or short circuits the recorded signal, reducing it by about half^{71,78}. For optogenetic experiments we additionally added silicon carbide powder (9 % wt, Tech-Met, Markham) to the grease to make it opaque to light and minimize light induced artifactual current in the silver-silver chloride recording wire during optogenetic activation of ChR2 (detailed below). Likewise, we covered our bath ground and recording return wires with a plastic shield to prevent stray light artifacts. The dorsal root recordings were amplified (2,000 times), high-pass filtered at 0.1 Hz to remove drift, low-pass filtered at 10 kHz, and sampled at 30 kHz (Axoscope 8; Axon Instruments/Molecular Devices, Burlingame, CA).

These grease gap recordings of PAD on sensory afferents reflect only the response of largest diameter axons in the dorsal root, mainly group I proprioceptive afferents, because of the following considerations. First, the largest axons in peripheral nerves have a nodal spacing of about 1 mm^{79,80}, and length constants λ_S are estimated to be similar, at about 1-2 times the nodal spacing⁸¹, Further, in our recordings we were only able to get the grease to within about 2 mm of the spinal cord. Thus, the centrally generated signal (PAD) is attenuated exponentially with distance x along the axon length in the bath (x = 2 mm). This is proportional to $\exp(-x/\lambda s)$ (see ⁷⁶), which is $1/e^2 = 0.11$ for x = 2 λs , as is approximately the case here. This makes a central PAD of about 4 mV appear as a ~0.4 mV potential on the root recording (DRP, 10 times smaller), as we previously reported 16. Furthermore, the nodal spacing and \(\lambda \) decrease linearly with smaller axon diameters \(^{76,79} \), making the voltages recorded on the smaller afferents contribute to much less of the compound root potential (halving the diameter attenuates PAD instead by 1/e⁴ or 0.012, which is 99% attenuation). Finally, unmyelinated sensory axons attenuate voltages over a much shorter distance than myelinated axons, since that membrane resistance (Rm) drops markedly without myelin and λs is proportional to $\sqrt{Rm/Ri}$ (where Ri is axial resistance; Stein 1980). Thus, any centrally generated change in potential in these small axons is unlikely to contribute to the recorded signal 2 mm away.

The composite EPSPs in many motoneurons were likewise recorded from the central cut end of ventral roots mounted in grease (grease gap), which has also previously been shown to yield reliable estimates of the EPSPs (though again attenuated by the distance from the motoneurons)⁸². The monosynaptic EPSPs were again identified as monosynaptic by their rapid onset (first component, ~1 ms after afferent

volley arrives in the ventral horn; see below), lack of variability in latency (< 1 ms jitter), persistence at high rates (10 Hz) and appearance in isolation at the threshold for DR stimulation (< 1.1xT), unlike polysynaptic reflexes which varying in latency, disappear at high rates, and mostly need stronger DR stimulation to activate.

Analysis of synaptic responses in sensory axons (PAD) and motoneurons (EPSPs).

When we recorded from sensory axons of an associated dorsal root (directly or via the dorsal roots) stimulation of an adjacent dorsal root (not containing the recorded axon; 0.1 ms, 1-3xT) evoked a characteristic large and long depolarization of the afferents, previously demonstrated to be mediated by GABAergic input onto the sensory axons ¹⁶. This depolarization is termed primary afferent depolarization (PAD). PAD occurs at a minimal latency of 4-5 ms following the afferent volley, consistent with its trisynaptic origin ^{14,16}, making it readily distinguishable from earlier events on the axon. PAD has a fast synaptic component evoked by a single DR stimulation (rising within 30 ms and decaying exponentially over < 100 ms; termed phasic PAD) and a slower longer lasting extrasynaptic component (starting at about 30 ms and lasting many seconds) that is enhanced by repeated DR stimulation (tonic PAD, especially with cutaneous stimulation) ¹⁶. We used this sensory activation of PAD or direct optogenetic activation of PAD to examine the action of GABA on sensory axon spike transmission to motoneurons, usually evoking phasic PAD about 10-60 ms prior to spikes or associated EPSPs on motoneurons (during phasic PAD), though we also examined longer lasting effects of tonic PAD evoked by repeated DR stimulation. Sometimes PAD is so large that it directly evokes spikes on the afferents, and these travel out the dorsal root, and thus they have been termed dorsal root reflexes (DRRs) ^{16,23}. We usually minimized these DRRs by keeping the DR stimulus that evokes PAD low (1.1 - 3.0 xT), though there were inevitably some DRRs, as they even occur in vivo in cats and humans 9,36,46.

When we recorded from motoneurons (directly or via ventral roots) stimulation of proprioceptive afferents in a dorsal root (0.1 ms, 1.1-1.5xT) evoked a monosynaptic EPSP, and associated monosynaptic reflex (MSR, spikes from EPSP). This EPSP is depressed by fast repetition (rate depended depression, RDD) ⁸³, and thus to study the EPSP we evoked it at long intervals (10 s, 0.1 Hz rate) where RDD was less. However, even with this slow repetition rate (0.1 Hz), at the start of testing the first EPSP was often not similar to the steady state EPSP after repeated testing. Thus, to avoid RDD we usually ran the 0.1 Hz EPSP testing continuously throughout the experiment, at least until a steady state response was reached (after 10 minutes). We then examined the action of activating (or inhibiting)

GABA_{axo} neurons on this steady state EPSP, by introducing light or sensory conditioning that activated these neurons at varying intervals (inter-stimulus intervals, ISIs) prior to each EPSP stimulation (control, GAD2//ChR2 mice and GAD2//Arch3 mice). We averaged the EPSP from ~10 trials (over 100 s) just before conditioning and then 10 trials during conditioning, and then computed the change in the peak size of the monosynaptic EPSP with conditioning from these averages. After conditioning was completed EPSP testing continued and any residual changes in the EPSP were computed from the 10 trials following conditioning (after-effect). Finally, EPSP testing continued over many minutes after which the original steady state EPSP was established. The background motoneuron potential, membrane resistance (Rm) and time constant just prior to the EPSP was also assessed before and after conditioning to examine whether there were any postsynaptic changes that might contribute to changes in the EPSP with conditioning. Along with the VR recordings, we simultaneously recorded PAD from DRs by similar averaging methods (10 trials of conditioning), to establish the relation of changes in EPSPs with associated sensory axon depolarization PAD.

Drugs and solutions

Two kinds of artificial cerebrospinal fluid (ACSF) were used in these experiments: a modified ACSF (mACSF) in the dissection chamber prior to recording and a normal ACSF (nACSF) in the recording chamber. The mACSF was composed of (in mM) 118 NaCl, 24 NaHCO3, 1.5 CaCl2, 3 KCl, 5 MgCl2, 1.4 NaH2PO4, 1.3 MgSO4, 25 D-glucose, and 1 kynurenic acid. Normal ACSF was composed of (in mM) 122 NaCl, 24 NaHCO3, 2.5 CaCl2, 3 KCl, 1 MgCl2, and 12 D-glucose. Both types of ACSF were saturated with 95% O2-5% CO2 and maintained at pH 7.4. The drugs sometimes added to the ACSF were APV (NMDA receptor antagonist), CNQX (AMPA antagonist), gabazine (GABAA antagonist), bicuculline (GABA_A, antagonist), L655708 (α5 GABA_A, antagonist), CGP55845 (GABA_B antagonist; all from Tocris, USA), 5-HT, kynurenic acid (all from Sigma-Aldrich, USA), and TTX (TTX-citrate; Toronto Research Chemicals, Toronto). Drugs were first dissolved as a 10 - 50 mM stock in water or DMSO before final dilution in ACSF. DMSO was necessary for dissolving gabazine, L655708, bicuculline and CGP55845, but was kept at a minimum (final DMSO concentration in ACSF < 0.04%), which by itself had no effect on reflexes or sensory axons in vehicle controls (not shown). L655708 was particularly difficult to dissolve and precipitated easily, especially after it had been exposed a few times to air; so immediately after purchase we dissolved the entire bottle and froze it at -40°C in single use 5 -20 μl aliquots, and upon use it was first diluted in 100 μl distilled water before dispersing it into ACSF.

Recording monosynaptic reflexes in awake mice and rats, and PAD activation.

Window implant over spinal cord. In GAD2//ChR2+ mice and control GAD2//ChR- mice a glass window was implanted over the exposed spinal cord to gain optical access to the sacrocaudal spinal cord, as described previously ⁶⁵. Briefly, mice were given Meloxicam (1 mg/kg, s.c.) and then anesthetized using ketamine hydrochloride (100 mg/kg, i.p.) and xylazine (10 mg/kg, i.p.). Using aseptic technique, a dorsal midline incision was made over the L2 to L5 vertebrae. Approximately 0.1 ml of Xylocaine (1%) was applied to the surgical area and then rinsed. The animals were suspended from a spinal-fork stereotaxic apparatus (Harvard Apparatus) and the muscles between the spinous and transverse processes were resected to expose the L2 to L5 vertebrae. The tips of modified staples were inserted along the lateral edge of the pedicles and below the lateral processes of L2 and L5, and glued in place using cyanoacrylate. A layer of cyanoacrylate was applied to all of the exposed tissue surrounding the exposed vertebrae followed by a layer of dental cement to cover the cyanoacrylate and to form a rigid ring around the exposed vertebrae. A modified paperclip was implanted in the layer of dental cement to serve as a holding point for surgery. A laminectomy was performed at L3 and L4 to expose the spinal cord caudal to the transection site. Approximately 0.1 ml of Xylocaine (1%) was applied directly to the spinal cord for 2-3 s, and then rinsed. A line of Kwik-Sil (World-Precision Instruments) was applied to the dura mater surface along the midline of the spinal cord and a glass window was immediately placed over the exposed spinal cord. The window was glued in place along the outer edges using cyanoacrylate followed by a ring of dental cement. Small nuts were mounted onto this ring to later bolt on a backpack to apply the laser light (on the day of experimentation). Saline (1 ml, s.c.) and buprenorphine (0.03 mg/kg, s.c.) was administered post-operatively, and analgesia was maintained with buprenorphine (0.03 mg/kg, s.c.) every 12 hours for two days. Experimentation started 1 week after the window implant when the mouse was fully recovered.

Percutaneous EMG wire implant and fibre optic cable attachment. On the day of experimentation, the mouse was briefly sedated with isoflurane (1.5 %) and fine stainless steel wires (AS 613, Cooner Wire, Chatsworth, USA) were percutaneously implanted in the tail for recording EMG and stimulating the caudal tail trunk nerve, as we previously detailed (wires de-insulated by 2 mm at their tip and inserted in the core of 27 gauge needle that was removed after insertion)⁶². A pair of wires separated by 8 mm were inserted at base of the tail for recording EMG the tail muscles, and second pair of wires was inserted nearby for bipolar activation of the caudal trunk nerve to evoke reflexes. A fifth ground wire was implanted between the EMG and stimulation wires. Following this a backpack was bolted into the nuts imbedded in the dental cement ring around the window. This backpack held and aligned a light fibre optic cable that was focused on the centre of the S3 – S4 sacral spinal cord. The Cooner wires

were secured to the skin with drops of cyanoacrylate and taped onto the backpack so that the mouse could not chew them. The isoflurane was removed, and the mouse quickly recovered from the anesthesia and was allowed to roam freely around an empty cage during recording, or was sometimes lightly restrained by hand or by a sling. The fibre optic cable was attached to a laser (447 nM, same above) and the Cooner wires attached to the same models of amplifiers and stimulators used for ex vivo monosynaptic testing detailed above.

MSR testing.

The monosynaptic reflex (MSR) was recorded in the tail EMG at ~6 ms latency after stimulating the caudal tail trunk nerve at a low intensity that just activated proprioceptive afferents (0.2 ms current pulses, 1.1 xT), usually near the threshold to activate motor axons and an associated M-wave (that arrived earlier). We studied the tail MSR reflex because our ex vivo recordings were made in the corresponding sacral spinal cord of adult mice and rats, which is the only portion of the spinal cord that survives whole ex vivo, due to its small diameter ⁶⁴. This reflex was verified to be of monosynaptic latency because it was the first reflex to arrive, had little onset jitter, and had the same latency as the F wave (not shown; the F wave is evoked by a strong stimulation of all motor axons, at 5xT, which causes a direct motoneuron response on the same axons, while the monosynaptic EPSP is blocked by collision at this intensity) ⁸⁴. The MSR also underwent rate dependent depression (RDD) with fast repeated stimulation and so was synaptic and not a direct muscle response (M-wave), which occurred earlier at sufficient intensity to recruit the motor axons (not shown).

Conditioning of the MSR by optogenetic activation of GABA_{axo} neurons. As with in vitro EPSP testing, the MSR was tested repeatedly at long 5-10 s intervals until a steady state MSR was achieved. Then testing continued but with a conditioning light pulse applied just prior to the MSR stimulation (40-120 ms), to examine the effect of PAD evoked during this time frame on sensory transmission to motoneurons. Background EMG just prior to MSR testing was assessed to estimate the postsynaptic activity on the motoneurons. The changes in MSR and background EMG with light were quantified by comparing the average response before and during the light application, computed from the mean rectified EMG at 6-11 ms after the nerve stimulation (MSR) and over 20 ms prior to the nerve stimulation (background just prior to the MSR, Bkg). Because awake mice spontaneously varied their EMG, we plotted the relation between the MSR and the background EMG, with as expected a positive linear relation between these two variables 85 , computed by fitting a regression line. In trials with conditioning light applied the same plot of EMG vs background EMG was made and a second

regression line computed. The change in the MSR with conditioning at a fixed matched background EMG level was then computed for each mouse by measuring the difference between the regression line responses at a fixed background EMG. This ruled out changes in MSRs being due to postsynaptic changes. Two background levels were assessed: rest (0%) and 30% of maximum EMG, expressed as a percentage of the control pre-conditioning MSR. The change in background EMG with light was computed by comparing the EMG just prior to the light application (over 20 ms prior) to the EMG just prior to the MSR (over 20 ms prior, Bkg), and expressed as a percentage of the maximum EMG.

Cutaneous conditioning of the MSR in rats. A similar examination of how PAD affected the MSR was performed in rats with percutaneous tail EMG recording. However, in this case PAD was evoked by a cutaneous conditioning stimulation of the tip of the tail (0.2 ms pulses, 3xT, 40-120 ms prior to MSR testing) using an additional pair of fine Cooner wires implanted at the tip of the tail (separated by 8 mm). In rats the MSR latency is later than in mice due to the larger peripheral conduction time, \sim 12 ms (as again confirmed by a similar latency to the F wave). This MSR was thus quantified by averaging rectified EMG over a 12-20 ms window. Also, to confirm the GABA_A receptor involvement in regulating the MSR, the antagonist L655708 was injected systemically (1 mg/kg i.p., dissolved in 50 μ l DMSO and diluted in 900 μ l saline). Again, the MSR was tested at matched background EMG levels before and after conditioning (or L655708 application) to rule out changes in postsynaptic inhibition.

Conditioning of the MSRs in humans

H-reflex as an estimate of the MSR. Participants were seated in a reclined, supine position on a padded table. The right leg was bent slightly to access the popliteal fossa and padded supports were added to facilitate complete relaxation of all leg muscles. A pair of Ag-AgCl electrodes (Kendall; Chicopee, MA, USA, 3.2 cm by 2.2 cm) was used to record surface EMG from the soleus muscle. The EMG signals were amplified by 1000 and band-pass filtered from 10 to 1000 Hz (Octopus, Bortec Technologies; Calgary, AB, Canada) and then digitized at a rate of 5000 Hz using Axoscope 10 hardware and software (Digidata 1400 Series, Axon Instruments, Union City, CA) ⁶². The tibial nerve was stimulated with an Ag-AgCl electrode (Kendall; Chicopee, MA, USA, 2.2 cm by 2.2 cm) in the popliteal fossa using a constant current stimulator (1 ms rectangular pulse, Digitimer DS7A, Hertfordshire, UK) to evoke an H-reflex in the soleus muscle, an estimate of the MSR ³⁷. Stimulation intensity was set to evoke a test (unconditioned) MSR below half maximum. MSRs recorded at rest were evoked every 5 seconds to minimize RDD ⁸⁶ and at least 20 test MSRs were evoked before conditioning to establish a steady baseline because the tibial nerve stimulation itself can presumably also activate spinal GABAergic

networks, as in rats. All MSR were recorded at rest, except when the motor unit firing probabilities were measured (see below).

Conditioning of the MSR. To condition the soleus MSR by cutaneous stimulation, the cutaneous medial branch of the deep peroneal (cDP) nerve was stimulated on the dorsal surface of the ankle using a bipolar arrangement (Ag-AgCl electrodes, Kendall; Chicopee, MA, USA, 2.2 cm by 2.2 cm), set at 1.0xT, where T is the threshold for cutaneous sensation. A brief burst (3 pulses, 200 Hz for 10 ms) of cDP stimuli was applied before evoking a MSR at various inter-stimulus intervals (ISIs; interval between tibial and cDP nerve stimuli) within the window expected for phasic PAD evoked by cutaneous stimuli, presented in random order at 0, 30, 60, 80, 100, 150 and 200 ms ISIs. Seven conditioned MSR at each ISI were measured consecutively and the average of these MSR (peak-to-peak) was used as an estimate of the conditioned MSR. This was compared to the average MSR without conditioning, computed from the 7 trials just prior to conditioning.

The cDP nerve was also stimulated with a 500 ms long train at 200 Hz to condition the MSR, and examine the effect of tonic PAD evoked by such long trains, as in rats. Following the application of at least 20 test MSRs (every 5 s), a single cDP train was applied 700 ms before the next MSR and following this the MSR continued to be evoked for another 90 to 120 s (time frame of tonic PAD). We also conditioned the soleus MSR with tibialis anterior (TA; antagonist muscle, flexor) tendon vibration (brief burst of 3 cycles of vibration at 200Hz) to preferentially activate Ia afferents, as has been done previously ³⁷.

Motor unit recording to examine postsynaptic actions of conditioning. Surface electrodes were used to record single motor units in the soleus muscle during low level contractions by placing electrodes on or near the tendon or laterally on the border of the muscle as detailed previously ⁸⁷. Alternatively, single motor unit activity from the soleus muscle was also recorded using a high density surface EMG electrode (OT Bioelettronica, Torino, Italy, Semi-disposable adhesive matrix, 64 electrodes, 5x13, 8 mm inter-electrode distance) with 3 ground straps wrapped around the ankle, above and below the knee. Signals were amplified (150 times), filtered (10 to 900 Hz) and digitized (16 bit at 5120 Hz) using the Quattrocento Bioelectrical signal amplifier and OTBioLab+ v.1.2.3.0 software (OT Bioelettronica, Torino, Italy). The EMG signal was decomposed into single motor units using custom MatLab software as per ⁸⁸. Intramuscular EMG was used to record MUs in one participant as detailed previously ⁸⁹ to verify single motor unit identification from surface EMG.

To determine if there were any postsynaptic effects from the conditioning stimulation on the motoneurons activated during the MSR, we examined whether the cDP nerve stimulation produced any changes in the tonic firing rate of single motor units, which gives a more accurate estimate of membrane potential changes in motoneurons compared to compound EMG. Single motor units were activated in the soleus muscle by the participant holding a small voluntary contraction of around 5% of maximum. Both auditory and visual feedback were used to keep the firing rates of the units steady while the conditioning cutaneous was applied every 3 to 5 seconds. The instantaneous firing frequency profiles from many stimulation trials were superimposed and time-locked to the onset of the conditioning stimulation to produce a peri-stimulus frequencygram (PSF, dots in Fig. 7biii), as previously detailed ^{89,90}. A mean firing profile resulting from the conditioning stimulation (PSF) was produced by averaging the frequency values in 20 ms bins across time post conditioning (thick lines in Fig. 7biii and 7ciii). To quantify if the conditioning stimulation changed the mean firing rate of the tonically firing motor units, the % change in the mean PSF rate was computed at the time when the H reflex was tested (vertical line in Fig. 7bii-iii).

Unitary EPSP estimates from PSF. To more directly examine if the facilitation in MSR resulted from changes in transmission in Ia afferents after cutaneous afferent conditioning, we measured changes in the firing probability of single motor units (MUs) during the brief MSR time-course (typically 30 to 45 ms post tibial nerve stimulation) with and without cDP nerve conditioning. Soleus MSRs were as usual evoked by stimulating the tibial nerve, but while the participant held a small voluntary plantarflexion to activate tonic firing of a few single motor units. The size of the MSR was set to just above reflex threshold (and when the M-wave was < 5% of maximum) so that single motor units at the time of the MSR could be distinguished from the compound potential from many units that make up the MSR ⁹¹. For a given trial run, test MSRs were evoked every 3-5 s for the first 100 s and then MSR testing continued for a further 100s, but with a cDP-conditioning train (50 ms, 200 Hz) applied 500 ms prior to each MSR testing stimulation. These repeated high frequency trains evoke a tonic PAD in rats that facilitates sensory conduction. A 500 ms ISI was used to ensure the firing rate of the motor unit returned to baseline before the MSR was evoked, and this is also outside of the range of phasic PAD. Approximately 40-50 usable test and conditioned firing rate profiles were produced for a single session where the motor units had a steady discharge rate before the cDP nerve stimulation. Sessions were repeated 3-6 times to obtain a sufficient number of frequency points to construct the PSF (~ 200 trials).

To estimate the EPSP profile, motor unit (MU) firing was again used to construct a PSF, as detailed above, but this time locked to the tibial nerve stimulation used to evoke the MSR, so that we could estimate the motoneuron behaviour during the MSR (EPSP). When more than one MU was visible in the recordings firing from these units (usually 2 – 3) were combined into a single PSF. Overall this gave about of 100 – 600 MU MSR test sweeps to generate each PSF. Firing frequency values were averaged in consecutive 20 ms bins to produce a mean PSF profile over time after tibial nerve stimulation, for both unconditioned and conditioned MSR reflex trials. The mean background firing rate within the 100 ms window immediately preceding the tibial stimulation was compared between the test and conditioned MSR trials to determine if the conditioning cDP nerve stimulation produced a change in firing rate, and thus post-synaptic effect, just before the conditioned MSR was evoked. For each PSF generated with or without conditioning, the probability that a motor unit discharged during the MSR window (30 to 45 ms after the TN stimulation) was measured as the number of discharges during the time of the MSR window divided by the total number of tibial nerve test stimuli. As an estimate of EPSP size, the mean firing rate during the MSR window was also measured (this was computed with smaller PSF bins of 0.5 ms during the MSR).

Temperature, latency and PAD considerations.

Large proprioceptive group Ia sensory afferents conduct in the peripheral tail nerve with a velocity of about 33 m/s (33 mm/ms) in mice 92 . Motor axons are similar, though slightly slower (30 m/s) 93 . Thus, in the awake mouse stimulation of Ia afferents in the mouse tail evokes spikes that take ~ 2 ms to conduct to the motoneurons in the spinal cord ~ 70 mm away. Following ~ 1 ms synaptic and spike initiation delay in motoneurons, spikes in the motor axons take a further ~ 2 ms to reach the muscles, after which the EMG is generated with a further 1 ms synaptic and spike initiation delay at the motor endplate to produce EMG. All told this gives a monosynaptic reflex latency of ~ 6 ms. The motor unit potentials within the EMG signal have a duration of about 3-5 ms, and thus we averaged rectified EMG over 6-11 ms to quantify the MSR. We have shown that similar considerations hold for the rat where tail nerve conduction velocities are similar, except the distance from the tail stimulation to the spinal cord is larger (150 mm), yielding a peripheral nerve conduction delay of ~ 10 ms and total MSR delay of ~ 12 ms 94 . In humans the MSR latency is dominated by the nerve conduction latency (50 -60 m/s) over a large distance (~ 800 mm), yielding MSR latencies of ~ 30 ms.

In our ex vivo whole adult spinal cord preparation the bath temperature was varied between 23 and 32° C. All data displayed is from $23 - 24^{\circ}$ C, though we confirmed the main results (facilitation of

sensory axon transmission to motoneuron by PAD) at 32°C. The Q10 for peripheral nerve conduction (ratio of conduction velocities with a 10 °C temperature rise) is about 1.3 95, yielding a conduction in dorsal roots of about 20 m/s at 23 – 24 °C, as we directly confirmed (not shown). Thus, when the DR is stimulated 20 mm from the cord the latency of spike arrival at the cord should be about 1 ms, which is consistent with the time of arrival of afferent volleys that were seen in the intracellular and extracellular recordings from sensory axons (e.g. Figs. 2b and 4e).

When we found that PAD evoked in sensory axons can prevent failure of spikes to propagate in the cord after DR stimulation, we worried that PAD somehow influenced the initiation of the spike by the dorsal root stimulation at the silver wire. However, we ruled this out by stimulating dorsal roots as far away from the spinal cord as possible (20 mm), where PAD has no effect, due to the exponential attenuation of its dorsal root potential with distance (see above), and found that PAD still facilitated sensory axon spike transmission to motoneurons. The added advantage of these long roots is that there is a clean 1 ms separation between the stimulus artifact and the afferent volley arriving at the spinal cord, allowing us to quantify small FPs and afferent volleys that are otherwise obscured by the artifact.

We did not consistently use high temperature ex vivo baths (32°C) because the VR and DR responses to activation of DRs or PAD neurons are irreversibly reduced by prolonged periods at these temperatures, suggesting that the increased metabolic load and insufficient oxygen penetration deep in the tissue damages the cord at these temperatures. Importantly, others have reported that in sensory axons PADevoked spikes (DRRs) are eliminated in a warm bath and argued that this means they are not present in vivo, and not able to evoked a motoneuron response ⁷, despite evidence to the contrary ^{9,46}. However, we find that PAD itself is reduced in a warm bath by the above irreversible damage, and it is thus not big enough to evoke spikes in sensory axons; thus, this does not tell us whether these spikes should be present or not in vivo. Actually, in vivo we sometimes observed that with optogenetic activation of GABA_{axo} neurons and associated PAD there was a direct excitation of the motoneurons (seen in the EMG) at the latency expected for PAD evoked spikes (not shown). However, this was also at the latency of the postsynaptic inhibition produced by this same optogenetic stimulation, which often masked the excitation (Fig. 6). In retrospect, examining the GABA_{axo} evoked motoneuron responses during optogenetic-evoked PAD (Fink et al.)^{7,17}, or sensory-evoked PAD^{17,18}, there is a small excitation riding on the postsynaptic IPSPs from the activation of there GABA_{axo} neurons. This is consistent with the PAD evoked spike activating the monosynaptic pathway, which inhibits subsequently tested monosynaptic responses by post activation depression (see Discussion).

The latency of a single synapse in our ex vivo preparation at 23 – 24°C was estimated from the difference between the time arrival of the sensory afferent volley at the motoneurons (terminal potential seen in intracellular and extracellular recordings) and the onset of the monosynaptic EPSP in motoneurons. This was consistently 1 – 1.2 ms (Fig. 5b and e). This is consistent with a Q10 of about 1.8 – 2.4 for synaptic transmission latency ^{96,97}, and 0.4 ms monsynaptic latency at body temperature ^{98,99}. Based on these considerations we confirm that the PAD evoked in sensory axons is monosynaptically produced by optogenetic activation of GABA_{axo} neurons with light, since it follows ~1 ms after the first spike evoked in GABAaxo neurons by light (Fig. 3a). This first spike in GABA_{axo} neurons itself takes 1 – 2 ms to arise and so the overall latency from light activation to PAD production can be 2 - 3 ms (Fig. 3f), as seen for IPSCs at this temperature in other preparations ¹⁰⁰. With DRs stimulation PAD arises with a minimally 4 – 5 ms latency, which is consistent with a trisynaptic activation of the sensory axon, after taking into account time for spikes to arise in the interneurons involved (Fig. 4a,e).

Immunohistochemisty.

Tissue fixation and sectioning. After sensory axons in mouse and rat spinal cords were injected with neurobiotin the spinal cord was left in the recording chamber in oxygenated nACSF for an additional 4 – 6 hr to allow time for diffusion of the label throughout the axon. Then the spinal cord was immersed in 4% paraformaldehyde (PFA; in phosphate buffer) for 20-22 hours at 4°C, cryoprotected in 30% sucrose in phosphate buffer for 24-48 hours. Alternatively, afferents were labelled genetically in VGLUT1^{Cre/+}; R26^{lsl-tdTom} mice, which were euthanized with Euthanyl (BimedaMTC; 700 mg/kg) and perfused intracardially with 10 ml of saline for 3 – 4 min, followed by 40 ml of 4% paraformaldehyde (PFA; in 0.1 M phosphate buffer at room temperature), over 15 min. Then spinal cords of these mice were post-fixed in PFA for 1 hr at 4°C, and then cryoprotected in 30% sucrose in phosphate buffer (~48 hrs). Following cryoprotection all cords were embedded in OCT (Sakura Finetek, Torrance, CA, USA), frozen at -60C with 2-methylbutane, cut on a cryostat NX70 (Fisher Scientific) in sagittal or transverse 25 μm sections, and mounted on slides. Slides were frozen until further use.

Immunolabelling. The tissue sections on slides were first rinsed with phosphate buffered saline (PBS, 100 mM, 10 min) and then again with PBS containing 0.3% Triton X-100 (PBS-TX, 10 min rinses used for all PBS-TX rinses). For the sodium channel antibody, we additionally incubated slides three times for 10 min each with a solution of 0.2% sodium borohydride (NaBH4, Fisher, S678-10) in PB, followed

by a PBS rinse (4x 5 min). Next, for all tissue, nonspecific binding was blocked with a 1 h incubation in PBS-TX with 10% normal goat serum (NGS; S-1000, Vector Laboratories, Burlingame, USA) or normal donkey serum (NDS; ab7475, Abcam, Cambridge, UK). Sections were then incubated for at least 20 hours at room temperature with a combination of the following primary antibodies in PBS-TX with 2% NGS or NDS: rabbit anti-α₅ GABA_A receptor subunit (1:200; TA338505, OriGene Tech., Rockville, USA), rabbit anti-α1 GABAA receptor subunit (1:300; 06-868, Sigma-Aldrich, St. Louis, USA), guinea pig anti-α2 GABAA receptor subunit (1:500; 224 104, Synaptic Systems, Goettingen, Germany), chicken anti-γ2 GABAA receptor subunit (1:500; 224 006, Synaptic Systems, Goettingen, Germany), rabbit anti-GABAB1 receptor subunit (1:500; 322 102, Synaptic Systems, Goettingen, Germany), mouse anti-Neurofilament 200 (NF200) (1:2000; N0142, Sigma-Aldrich, St. Louis, USA), guinea pig anti-VGLUT1 (1:1000; AB5905, Sigma-Aldrich, St. Louis, USA), rabbit anti-Caspr (1:500; ab34151, Abcam, Cambridge, UK), mouse anti-Caspr (1:500; K65/35, NeuroMab, Davis, USA), chicken anti-Myelin Basic Protein (MBP) (1:200; ab106583, Abcam, Cambridge, UK), chicken anti-VGAT (1:500; 131 006, Synaptic Systems, Goettingen, Germany), rabbit anti-VGAT (1:500; AB5062P, Sigma-Aldrich, St. Louis, USA), rabbit anti-EYFP (1:500; orb256069, Biorbyt, Riverside, UK), goat anti-RFP (1:500; orb334992, Biorbyt, Riverside, UK), rabbit anti-RFP (1:500; PM005, MBL International, Woburn, USA), rabbit anti-GFP (1:500, A11122, ThermoFisher Scientific, Waltham, USA), and mouse anti-Pan Sodium Channel (1:500; S8809, Sigma-Aldrich, St. Louis, USA). The latter is a pan-sodium antibody, labelling an intracellular peptide sequence common to all known vertebrate sodium channels. Genetically expressed EYFP, tdTom (RFP) and GFP were amplified with the above antibodies, rather than rely on the endogenous fluorescence. When anti-mouse antibodies were applied in mice tissue, the M.O.M (Mouse on Mouse) immunodetection kit was used (M.O.M; BMK-2201, Vector Laboratories, Burlingame, USA) prior to applying antibodies. This process included 1h incubation with a mouse Ig blocking reagent. Primary and secondary antibody solutions were diluted in a specific M.O.M diluent.

The following day, tissue was rinsed with PBS-TX (3x 10 min) and incubated with fluorescent secondary antibodies. The secondary antibodies used included: goat anti-rabbit Alexa Fluor 555 (1:200; A32732, ThermoFisher Scientific, Waltham, USA), goat anti-rabbit Alexa Fluor 647 (1:500, ab150079, Abcam, Cambridge, UK), goat anti-rabbit Pacific orange (1:500; P31584, ThermoFisher Scientific, Waltham, USA), goat anti-mouse Alexa Fluor 647 (1:500; A21235, ThermoFisher Scientific, Waltham, USA), goat anti-mouse Alexa Fluor 488 (1:500; A11001, ThermoFisher Scientific, Waltham, USA), goat anti-mouse Alexa Fluor 555 (1:500; A28180, ThermoFisher Scientific, Waltham, USA), goat anti-mouse Alexa Fluor 555 (1:500; A28180, ThermoFisher Scientific, Waltham, USA), goat anti-

guinea pig Alexa Fluor 647 (1:500; A21450, ThermoFisher Scientific, Waltham, USA), goat antichicken Alexa Fluor 405 (1:200; ab175674, Abcam, Cambridge, UK), goat anti-chicken Alexa Fluor 647 (1:500; A21449, ThermoFisher Scientific, Waltham, USA), donkey anti-goat Alexa Fluor 555 (1:500; ab150130, Abcam, Cambridge, UK), donkey anti-rabbit Alexa Fluor 488 (1:500; A21206, ThermoFisher Scientific, Waltham, USA), Streptavidin-conjugated Alexa Fluor 488 (1:200; 016-540-084, Jackson immunoResearch, West Grove, USA) or Streptavidin-conjugated Cyanine Cy5 (1:200; 016-170-084, Jackson immunoResearch, West Grove, USA) in PBS-TX with 2% NGS or NDS, applied on slides for 2 h at room temperature. The latter streptavidin antibodies were used to label neurobiotin filled afferents. After rinsing with PBS-TX (2 times x 10 min/each) and PBS (2 times x 10 min/each), the slides were covered with Fluoromount-G (00-4958-02, ThermoFisher Scientific, Waltham, USA) and coverslips (#1.5, 0.175 mm, 12-544-E; Fisher Scientific, Pittsburg, USA).

Standard negative controls in which the primary antibody was either 1) omitted or 2) blocked with its antigen (quenching) were used to confirm the selectivity of the antibody staining, and no specific staining was observed in these controls. Most antibodies had been previously tested with quenching for selectivity, as detailed in the manufacture's literature and other publications ¹⁶, but we verified this for the GABA receptors with quenching. For antibody quenching, the peptides used to generate the antibodies, including anti-α5 GABAA receptor subunit (AAP34984, Aviva Systems Biology, San Diego, USA), anti-α1 GABAA receptor subunit (224-2P, Synaptic Systems, Goettingen, Germany) and anti-γ2 GABAA receptor subunit (224-1P, Synaptic Systems, Goettingen, Germany), were mixed with the antibodies at a 10:1 ratio and incubated for 20 h and 4°C. This mixture was then used instead of the antibody in the above staining procedure.

Confocal and epifluorescence microscopy

Image acquisition was performed by confocal (Leica TCS SP8 Confocal System) and epifluorescence (Leica DM 6000 B) microscopy for high magnification 3D reconstruction and low magnification imaging, respectively. All the confocal images were taken with a 63x (1.4 NA) oil immersion objective lens and 0.1 µm optical sections that were collected into a z-stack over 10–20 µm. Excitation and recording wavelengths were set to optimize the selectivity of imaging the fluorescent secondary antibodies. The same parameters of laser intensity, gain and pinhole size was used to take pictures for each animal, including the negative controls. Complete sagittal sections were imaged with an epifluorescence 10x objective lens using the Tilescan option in Leica Application Suite X software (Leica Microsystems CMS GmbH, Germany). Sequential low power images were used to reconstruct

the afferent extent over the whole spinal cord, using CorelDraw (Ottawa, Canada), and to identify locations where confocal images were taken.

3D reconstruction of afferents and localization of GABA receptors.

The fluorescently labelled afferents (neurobiotin, tdTom), GABA receptors, VGLUT1, VGAT, NF200, Caspr, MBP and sodium channels were analyzed by 3D confocal reconstruction software in the Leica Application Suite X (Leica Microsystems CMS GmbH) ¹⁶. To be very conservative in avoiding nonspecific antibody staining, a threshold was set for each fluorescence signal at a minimal level where no background staining was observed in control tissue with the primary antibody omitted, less 10%. Signals above this threshold were rendered in 3D for each antibody. Any GABA receptor, Caspr or Nav expression within the volume of the neurobiotin filled axon (binary mask set by threshold) was labelled yellow, pink and white respectively (Fig. 1), and the density within the afferents quantified using the same Leica software. Receptor densities were measured for all orders of branch sizes (1st, 2nd, 3rd etc.; see below), for both branches dorsal to the central canal (dorsal) and ventral to the central canal (ventral). Nodes were identified with dense bands of Caspr or Na channel labelling (and lack of MBP). Branch points were also identified. We also examined raw image stacks of the neurobiotin afferents and receptors, to confirm that the automatically 3D reconstructed and identified receptors labelled within the afferent (yellow) corresponded to manually identified receptors colocalized with neurobiotin (Fig. 1). This was repeated for a minimum of 10 examples for each condition, and in all cases the 3D identified and manually identified receptors and channels were identical. Many receptors and channels lay outside the afferent, and near the afferent these were difficult to manually identify without the 3D reconstruction software, making the 3D reconstruction the only practical method to fully quantify the receptors over the entire afferent. We also optimized the reconstruction of the neurobiotin filled afferents following the methods of Fenrich¹⁰¹, including brightening and widening the image edges slightly (1 voxel, 0.001 µm³) when necessary to join broken segments of the afferent in the final afferent reconstruction, to account for the a priori knowledge that afferents are continuous and neurobiotin signals tend to be weaker at the membrane (image edges) and in fine processes.

GABA receptors usually occurred in the axons in distinct clusters. The distances between these receptor clusters and nodes or branch points was measured and average distances computed, from high power confocal images evenly sampled across the axon arbour (employing ~100 receptor clusters from ~10 images per labelled Ia afferent). The average distance between the receptor clusters and the nearest axon terminals on the motoneurons was also computed, but this was complicated by the very large distances

often involved, forcing us to compute the distances from low power images and relate these to the high power images of receptors sampled relatively evenly along the axon arbour. For this distance calculation, to avoid sampling bias in the high power images, we only admitted images from branch segments (1st, 2nd and 3rd order, detailed below) that had a receptor density within one standard deviation of the mean density in branch types with the highest density (1st order ventral branches for GABA_A receptors and 3rd order ventral terminal branches for GABA_B receptors; i.e. images from branches with density above the dashed confidence interval lines in Fig. 1g were included); this eliminated very large distances being included from branch segments with relatively insignificant receptor densities. We also confirmed these calculations by computing the weighted sum of all the receptor distances weighted by the sum of the receptor density for each branch type (and divided by the sum of all receptor densities), which further eliminated sampling bias. This gave similar average distance results to the above simpler analysis (not shown).

Sensory axon branch order terminology. The branches of proprioceptive axons were denoted as follows: dorsal column branches, 1st order branches that arose of the dorsal column and project toward the motoneurons, 2nd order branches that arose from the 1st order branches, and 3rd order branches that arose from the 2nd order branches. Higher order branches occasionally arose from the 3rd order branches, but these were collectively denoted 3rd order branches. First and second order branches were myelinated with large dense clusters of sodium channels at the nodes in the myelin gaps, which were characteristically widely spaced. As the second order branches thinned near the transition to 3rd order branches, they became unmyelinated, and at this point sodium channel clusters were smaller and more closely spaced (~6 µm apart, not shown). These thinned branches gave off 3rd order (and higher) unmyelinated terminal branches with chains of characteristic terminal boutons. The 1st order branches gave off 2nd order branches along most of their length as they traversed the cord from the dorsal columns to the motoneurons, but we separately quantified 1st, 2nd and 3rd order branches in the dorsal and ventral horn.

Node identification. Nodes in myelinated axon segments nodes were identified either directly via direct Na channel clusters and paranodal Caspr, or indirectly by their characteristic paranodal taper. That is, in the paranodal region the neurobiotin filled portion of the axon tapered to a smaller diameter, likely because the Caspr and presumably other proteins displaced the cytoplasmic neurobiotin, which also made the intracellular neurobiotin label less dense (Fig. 1c, black regions in taper). Regardless of the details, this taper made nodes readily identifiable. This taper forces the axial current densities to

increase at the nodes, presumably assisting spike initiation, and consistent with previous reconstructions of myelinated proprioceptive afferents ¹⁰².

Computer simulations

All computer models and simulations were implemented in NEURON ver7.5 103. The geometry and myelination pattern of the model were extracted from a previous study that used serial-section electron microscopy to generate about 15,000 photomicrographs to reconstruct a large myelinated proprioceptive Ia afferent collateral in the cat (Nicol and Walmsley, 1991). This structure was used in a prior modeling study ³⁴. Four classes of segment were defined in the model: myelinated internodes, nodes, unmyelinated bridges, and terminal boutons. Data from 18 of the 83 segments were missing from the original study. The missing data were estimated using mean values of the same segment class. The cable properties of the model were determined from diameter-dependent equations previously used for models of myelinated axons 104 and included explicit representation of myelinated segments using the double cable approach 104-106. Hodgkin-Huxley style models of voltage gated sodium (transient and persistent) and potassium channels were adopted from a previous study, at 37°C104. All three voltagegated conductances were colocalized to unmyelinated nodes and segments throughout the modelled axon collateral. The density of sodium and potassium conductances was adjusted to match the size and shape of experimentally recorded action potentials. To be conservative, sodium channels were placed at each node and bouton ($gNa = 1 \text{ S/cm}^2$), even though bouton immunolabelling for these channels was not common in our terminal bouton imaging (Fig. 1), since disperse weak sodium channel labelling may have been missed. Removing these bouton sodium channels did not qualitatively change our computer simulation results (not shown). Current clamp stimulation was applied to the middle of the first myelinated segment (pulse width 0.1 ms, amplitude 2 nA; near dorsal root) to initiate propagating action potentials in the model. Voltage at multiple sites of interest along the collateral was measured to assess propagation of action and graded potentials through branch points. Transient chloride conductance (i.e. GABA_A receptors) was modeled using a double-exponential point process (Eq. 1); parameters were manually fit to experimental data. GABAA receptors were localized to nodes at branch points to match experimental data. The amplitude and time course of the modeled PAD (also termed PAD) was measured from the first myelinated internode segment, similar to the location of our intraaxonal recordings.

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g(t) = g_max * \beta * (e^((-t)/\tau_decay) - e^((-t)/\tau_rise)), where \beta = 1/((e^((-\gamma)/\tau_decay) - e^((-\gamma)/\tau_rise))), and \gamma = (\tau_rise * \tau_decay)/(\tau_decay - \tau_rise) * \log(\tau_decay/\tau_rise) (Eq. 1)
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The parameters at all synapses were the same: time constant of rise (τ_{rise}) = 6ms, time constant of decay (τ_{decay}) = 50ms, default maximum conductance (g_{max}) = 1.5nS (varied depending on simulation, see figure legends), and chloride reversal potential (E_{Cl}) = -25mV (i.e. 55 mV positive to the resting potential to match our experimental data) ¹⁶. Space constants (λ_s) were computed for each segment of the afferent, from subthreshold current injections (100 ms) on the distal end of each branch segment and fitting an exponential decay (with space constant λ_s) to the passive depolarization along its length, and then repeating this with current injected in the proximal end to get a second λS , and finally averaging these two space constants.

QUANTIFICATION AND STATISTICAL ANALYSIS

Data were analyzed in Clampfit 8.0 (Axon Instruments, USA) and Sigmaplot (Systat Software, USA). A Student's t-test or ANOVA (as appropriate) was used to test for statistical differences between variables, with a significance level of P < 0.05. Power of tests was computed with $\alpha = 0.05$ to design experiments. A Kolmogorov-Smirnov test for normality was applied to the data set, with a P < 0.05 level set for significance. Most data sets were found to be normally distributed, as is required for a t-test. For those that were not normal a Wilcoxon Signed Rank Test was instead used with P < 0.05. Categorical data was instead analyzed using Chi-squared tests, with Yate's continuity correction used for 2×2 contingency tables and again significant difference set at P < 0.05. Data are plotted as bar graphs of mean \pm standard deviation (SD, error bar representing variability) or as box plots representing the interquartile range and median and error bars extend to the most extreme data point within 1.5 times the interquartile range (mean also shown as thick lines in boxes).

Data availability

All data are available in the manuscript or the supplementary materials. Raw data are available upon request to the corresponding authors. This study did not generate data sets or new unique reagents.

Code availability

The computer code used to perform the axon simulations (Extended Date Fig. 4) are publicly available on the github repository: https://github.com/kelvinejones/noah-axon.git

Acknowledgements

We thank Leo Sanelli, Jennifer Duchcherer, Babak Afsharipour and Christopher K. Thompson for technical assistance, and Shawn Hochman, CJ Heckman, FJ Alvarez and Tia Bennett for discussions and editing the manuscript. VGLUT1^{Cre} mice were kindly donated by Dr. Francisco J. Alvarez. This research was supported by the Canadian Institutes of Health Research (MOP 14697 and PJT 165823 D.J.B.) and the US National Institutes of Health (NIH, R01NS47567; D.J.B. and K.F.).

Author information

Contributions. K.H, and A.M.L-O. designed the study, carried out the animal experiments and analyzed data. K.M. and M.A.G. designed and performed the human experiments. N.P. and K.E.J. designed and performed the computer simulations. S.L., S.B., A.M., M.J.S. and R.S. assisted with animal electrophysiology. K.F. and A.M.L-O provided confocal microscopy. K.K.F, S.L., D.A.R. and K.H. developed the transgenic mice and performed the optogenetic experiments. Y.L. and D.J.B. conceived and designed the study, carried out experiments and analyzed data. D.J.B, K.H., and Y.L. wrote the paper, with editing from other authors. These authors contributed equally: Krishnapriya (Veni) Hari and Ana M. Lucas-Osma. These authors jointly supervised this work as senior authors: Yaqing Li, Keith K. Fenrich and David J. Bennett.

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Ethical declarations

All authors declare no competing interests.

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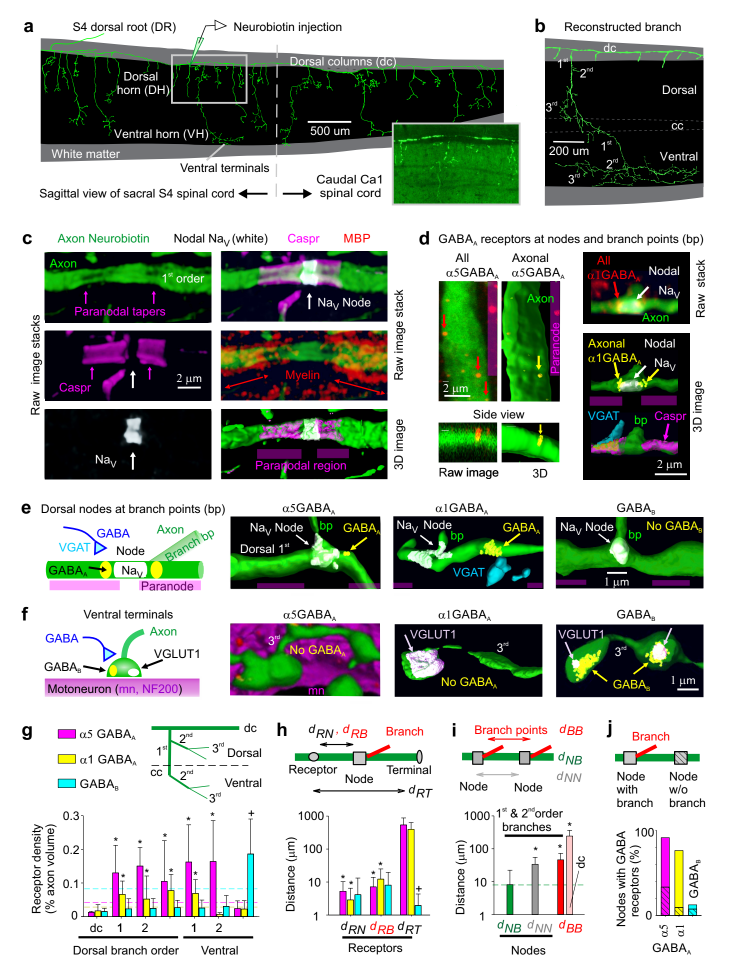


Fig. 1 | Nodal GABA_A and terminal GABA_B receptors in rats. a,b, Neurobiotin filled proprioceptive Ia axon in the sacrocaudal rat spinal cord (S4 and Ca1), reconstructed from fluorescent images (inset), with branching order indicated. Some ventral branches truncated in (a), but full arbour shown in (b). Central canal: cc. Dorsal columns: dc. Dorsal and ventral horns: DH and VH. c, Node on axon branch in DH immunolabelled for sodium channels (Nay), paranodal Caspr and myelin (MBP), with the paranodal taper indicated, and co-labelling within the axon rendered in 3D (bottom). 1st order branch in DH. d-f, α5 GABAA, α1 GABAA and GABAB receptor immunolabelling (yellow) in 3D reconstructed axon branches in DH (also raw images in \mathbf{d} , for $\alpha 5$ and $\alpha 1$), with nodes identified by Nav and paranodal taper, GABA contacts by vesicular inhibitory amino acid transporter (VGAT, e) labelling, and terminals by vesicular glutamate transporter 1 (VGLUT1, f) or nearby NF200 (on motoneurons in VH). 1st to 2nd order branch points (bp) in DH. g, Receptor densities on axon branches of varying order. Mean $\pm SD$ plotted. Dashed lines: lower confidence interval, 1 SD below mean maximum density. *significantly more than ventral terminal (3rd order) receptor density, + ventral terminal receptor density significantly more than 1^{st} and 2^{nd} order branch densities, P < 0.05, n = 10 - 17 branches per condition. **h**, Distances between GABA receptor clusters and nodes (d_{RN}) , branch points (d_{RB}) or terminals (d_{RT}) . *significantly less than d_{RT} . + significantly less than d_{RN} and d_{RB} , P < 0.05, n = 80 - 110 clusters per condition. i, Distances between branch points (d_{BB}), nodes (Nav clusters, d_{NN}), and nearest branch to node (d_{NB}). *significantly larger than d_{NB} , P < 0.05, n = 60 - 95 per condition. j, Proportion of nodes with GABA receptors, with and without (hashed) nearby branch points.

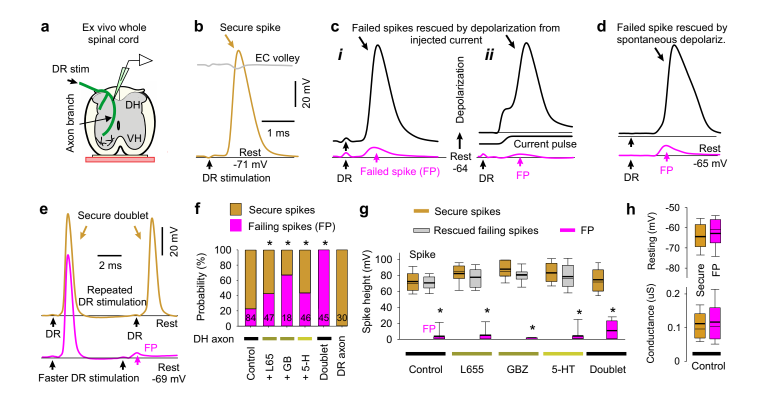


Fig. 2 | Spike failure. a, Recording from ex vivo whole adult rat spinal cord. b-d, Intracellular recordings from proprioceptive axon branches in the dorsal horn (DH), with dorsal root (DR) stimulation (1.1x threshold T, 0.1 ms) evoking a spike in some branches (secure, b) and only a failed spike in others (failure potential, FPs; c, d), but depolarization restoring full spikes (black, c and d). Averages of 10 trials at 3 s intervals. Resting potential: thin line. EC: extracellular afferent volley. Axons from S4 sacral DR. e, Fast repeated DR stimulation induced failure in secure spikes (not failing on first stimulation, 1.1xT). Threshold interval (longest) for failure (FP, pink), and just prior to threshold (gold). f, Proportions of DH axon branches (or DR axons) failing to spike with DR stimulation under control resting conditions and with L655708 (0.3 μM), gabazine (50 μM; GBZ), 5-HT (10 μM) or fast repetition (doublet, e). *significantly more than control, χ-squared test, P < 0.05, n indicated in bars. g, Summary of spike and FP heights in secure and failing branches. Box plots: median, thin line; mean, thick line; interquartile range IQR, box; extreme data points within 1.5 x IQR of box, error bars. *significantly smaller than secure spike, P < 0.05, for spikes in (f). h, Resting membrane potential and conductance for secure and failing branches, not significantly different, P > 0.05, for control axons in (f).

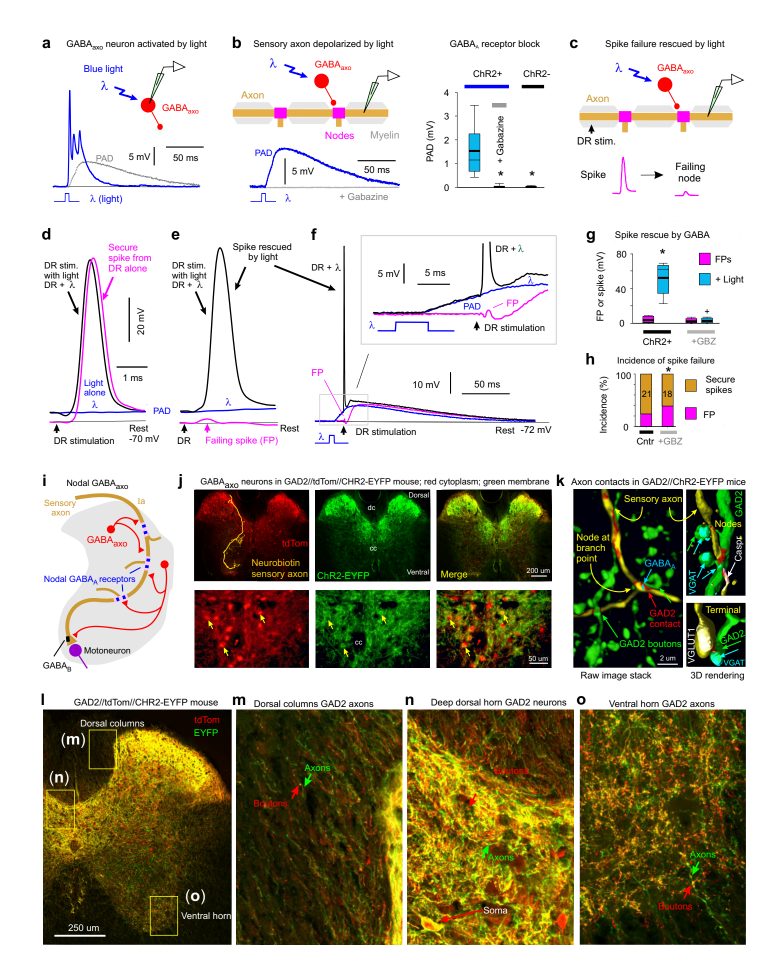


Fig. 3 | Nodal facilitation by GABA_{axo} neurons. a, Intracellular recording from GABA_{axo} neuron in ex vivo spinal cord of GAD2//ChR2-EYFP mouse, with ChR2 activated with a light pulse (5 ms, λ = 447nm laser, 0.7 mW/mm², 1.5x light threshold, T) causing a long depolarization and asynchronous spiking; isolated in 50 μ M gabazine (n = 5/5 similar). Average of 10 trials at 0.3 Hz, blue. PAD from (b), grey. Resting at -61mV. b, Intracellular recording from proprioceptive axon branch (in DH, sacral S3) with light pulse (1.5xT) producing a long depolarization (PAD, n = 14). Average of 10 trials at 0.3 Hz. *significantly less with gabazine or omitting ChR2 (control mice), n = 12 each. Resting at -71 mV. **c-g**, DR stimulation at rest (1.1xT) evoked a secure spike in some axon branches (**d**) and not others (**e**, **f**, FPs; DH S3 axons). Light evoked PAD (λ , 1.5xT, 10 ms prior) rescued failed spikes (e, f) and sped up conduction in secure spikes (d). Box plots of FPs and spikes (g); *significant increase with light, n =10; + significant reduction in light effect with 50 μ M gabazine, n = 7. h, Incidence of branches with failed DR-evoked spikes. *significant change with gabazine, γ -squared test, P < 0.05, n indicated. i-o, GABAaxo neurons imaged in S3 sacral spinal cord of GAD2//ChR2-EYFP//tdTom (i, l-o; green/red, merge yellow) or GAD2//ChR2-EYFP (k, green, dorsal horn) mice. Innervation of 3D reconstructed neurobiotin filled sensory axons (gold in j and k, as in Fig. 1) by GABA_{axo} neurons (green; axon contacts labelled red in k) in dorsal horn. Nodes identified by Caspr and paranodal taper, sensory terminals by VGLUT1, GABA_{axo} terminals by VGAT, and axonal GABA_A receptors by the α5GABA_A subunit. ChR2-EYFP is mainly expressed on plasma membranes 107, whereas tdTom is cytoplasmic, and so tdTom rings EYFP labelling, especially evident on the soma and boutons (j, l-o). Regions in (l) expanded in (m-o).

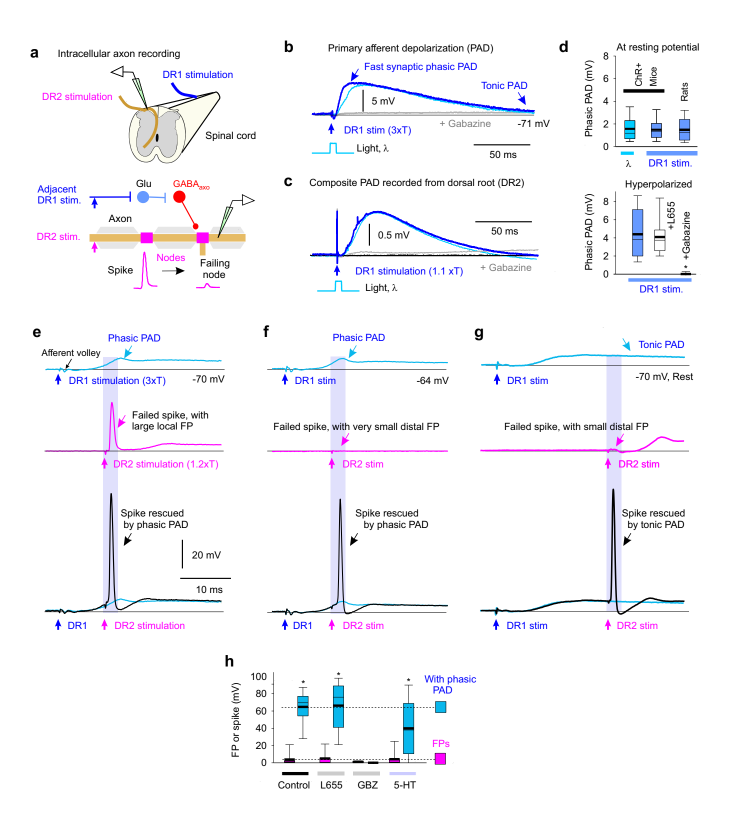


Fig. 4 | Sensory driven nodal facilitation. a, Experimental setup to indirectly activate GABAaxo neurons by DR stimulation (DR1), via a trisynaptic circuits detailed in Extended Data Figs. 5-6, in ex vivo spinal cords of rats or GAD2//ChR2-EYFP mice. b-c, Depolarization of a proprioceptive axon branch (b, intracellular in sacral S3 DH) or multiple axons in a DR (c, grease gap recording; sacral S3 DR; DR2) from stimulating the adjacent S4 DR (1.1xT, 0.1 ms pulse; DR1) or applying a light pulse to activate GABA_{axo} neurons (5 ms, 447nm, 0.7 mW/mm², as in Fig. 3b), both blocked by gabazine (50 uM; PAD), in GAD2//ChR2 mouse. Thin line resting potential. d, Summary box plots of peak phasic PAD evoked in axons by adjacent DR stimulation (DR1) or light, at rest (top, rats and mice, n = 14each) and with hyperpolarization (-10 mV, bottom, same rats), and effects of applied gabazine (50 µM; n = 14 rats) or L655708 (0.3 μ M; n = 8 rats). *significant difference from pre-drug (blue, lower plot), P < 0.05. e-g, DR axon branches (sacral S3 DH) exhibiting spike failure (FPs, magenta) following stimulating their DR (S3 DR, 1.1xT, 0.1 ms; DR2) in rats at rest. Spikes rescued by PAD evoked by prior conditioning of adjacent DR (S4 or contralateral S3 DR, at 3xT; DR1). Rescue occurred with fast synaptic depolarizations (phasic PAD; e-f) and tonic depolarizations (tonic PAD, g), both for local FPs (large, e) or distal FPs (small, f-g). h, FP or spike heights before and during DR evoked phasic PAD (n = 10) as in e-f, and actions of L655708 ($n = 9, 0.3 \mu M$), gabazine ($n = 14, 50 \mu M$) and 5-HT (n = 8, 10 μ M) in rats. *significant increase in spike with PAD, P < 0.05.

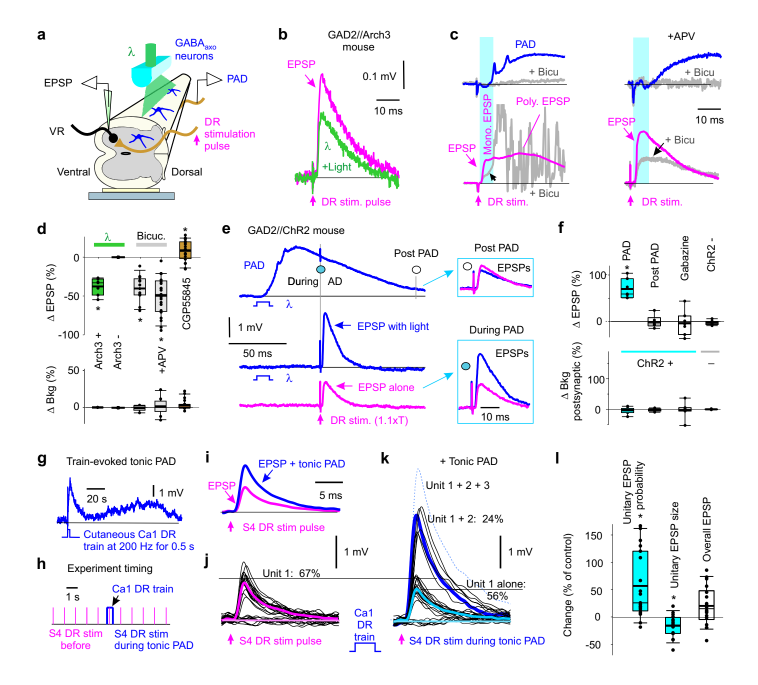


Fig. 5 | Facilitation of monosynaptic sensory transmission by GABA. a, Ex vivo recording from motoneurons while illuminating GABA_{axo} neurons with light λ . **b-d**, Composite monosynaptic EPSP in motoneurons (recorded in sacral S4 VR) evoked by a DR stimulation pulse alone (S4 DR, 0.1 ms, 1.1xT, magenta). Actions of optogenetic silencing GABA_{axo} neurons with light (a-b, 532nm, 5 mW/mm², 80 ms, in GAD2//Arch3 mice, n = 7), blocking GABA_A receptors (c, with bicuculline or gabazine, 50 μ M, with and without NMDA antagonist APV, 50 μ M; n = 23 and 13 in rats and mice combined), or blocking GABA_B receptors (**d**, CGP55845, 0.3 μ M, n = 10 rats and 10 mice). PAD shown for reference, recorded on S3 DR (c, top). Box plots of changes in EPSP and background postsynaptic activity (Bkg, over 10 ms prior to EPSP) with light or drugs, and with Arch3+ and Arch3-(n = 5) mice (d). * significant change Δ , P < 0.05. GAD2//Arch3 mice are VGAT+, not shown. e-f, Composite EPSP (evoked in S4 motoneurons, as in a-b) before, during and post PAD (recorded simultaneously on S3 DR) evoked by light activation of GABAaxo neurons (10 ms, 1.1xT, 447nm, 0.5 mW/mm², 60 ms and 140 ms pre EPSP, ISI) in GAD2//ChR2 mice. Box plots of changes in EPSP and Bkg (10 ms prior) with light in ChR2⁺ mice without and with gabazine (50 µM, during PAD), and in ChR2⁻ mice (60 ms ISI). * significant change, P < 0.05, n = 7 each. g, Tonic PAD (L655905 sensitive) recorded in sacral S4 proprioceptive axon in response to 0.5 s, 200 Hz DR stimulation train applied to the largely cutaneous Ca1 DR of caudal cord (3xT, DR2) in rat. h-i, Average EPSP in S4 motoneuron (intracellular recording, EPSP evoked by S4 DR stimulation at 3 s intervals used for average; DR1) before and during tonic PAD evoked by the brief DR train of g, at matched postsynaptic potentials (Bkg). **i-k**, Individual trials used to make EPSP averages in (i) (at 1 s intervals, h), with large all or nothing unitary EPSPs (thick lines unitary averages; dotted single occurrence of Unit 3). Lowpass filtered at 3 kHz. I, Changes in unitary EPSP probability and size, and overall EPSP with tonic PAD. Box plots. * significant change, P < 0.05, n = 18.

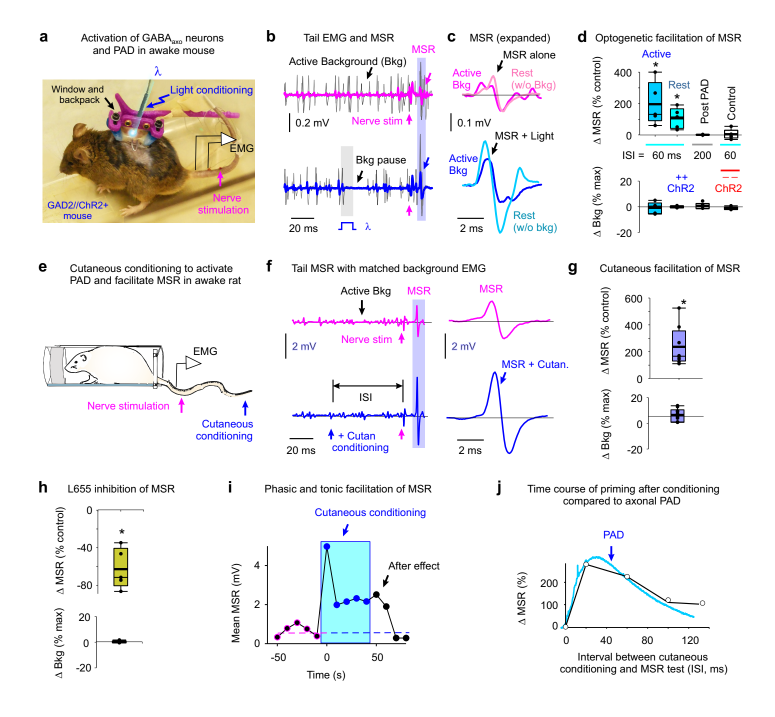


Fig. 6 | Facilitation of reflexes in awake animals. a, Recording tail muscle EMG and evoking monosynaptic reflexes (MSR) with tail nerve stimulation (1.1xT, 0.2 Hz), while activating GABA_{axo} neurons (PAD) with light ($\lambda = 447$ nm, 10 ms pulse, 1.5xT, 5 mW/mm²) in GAD2//ChR2 mice. **b-c**, Effect of light pulse λ on active background EMG (Active Bkg condition in **b**) and the MSR evoked 60 ms later, the latter expanded in (c). MSR tested with (b,c, Active Bkg, 30% max) and without (c, Rest) background EMG. Thin lines in (b) are individual trial examples at 10 s intervals (0.1 Hz); thick lines: averages. d, Changes in MSR with light activation of GABAaxo neurons at matched postsynaptic background (Bkg) (over 20 ms prior to MSR; lack of change in Bkg). Measured in active and resting (no Bkg) states, in ChR2+ and ChR2- mice (rest only), and during (60 ms ISI) and post PAD (200ms ISI at rest only). ISI: interstimulus interval. Box plots. * significant change, P < 0.05, n = 5 mice each. e-g, MSR recorded as in (b-d), but in rat and with PAD instead activated with cutaneous conditioning (tip of tail, 0.2 ms, 2xT, 60 ms prior, 0.1 Hz repetition), at matched active Bkg EMG. * significant change, P < 0.05, n = 8 rats. h, Decrease in MSR with L655935 (1 mg/kg i.p.) at matched Bkg EMG. Box plot. * significant change, P < 0.05, n = 5 rats. i, Typical MSR amplitude before, during and after conditioning as in (e) with after effect. i, Typical change in MSR with cutaneous conditioning as in (e) when the ISI is increased, compared to PAD (from Fig. 4). (i,j) similar results in n = 5/5 rats.

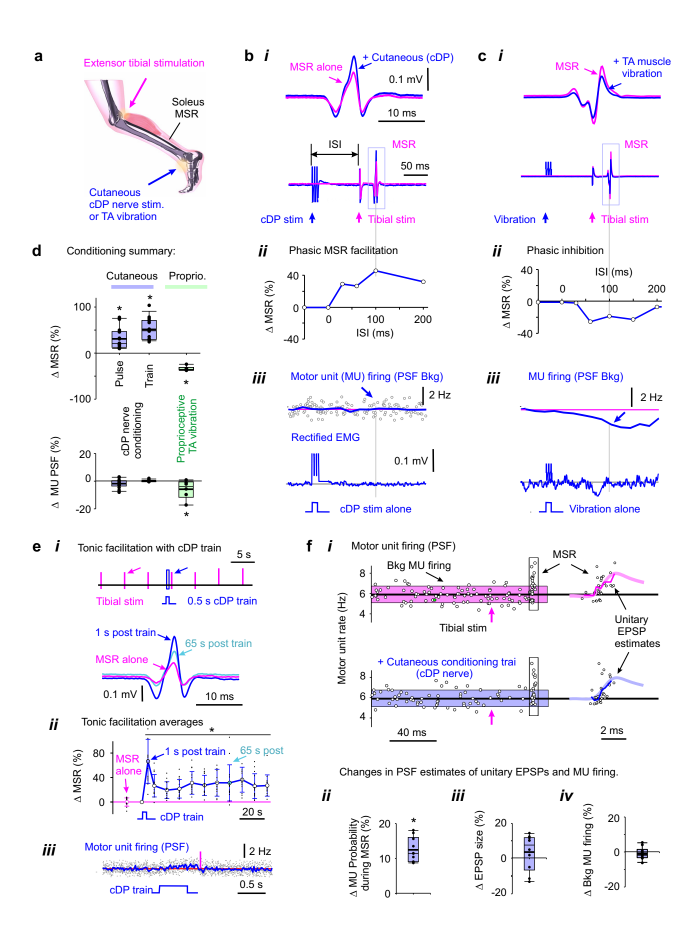


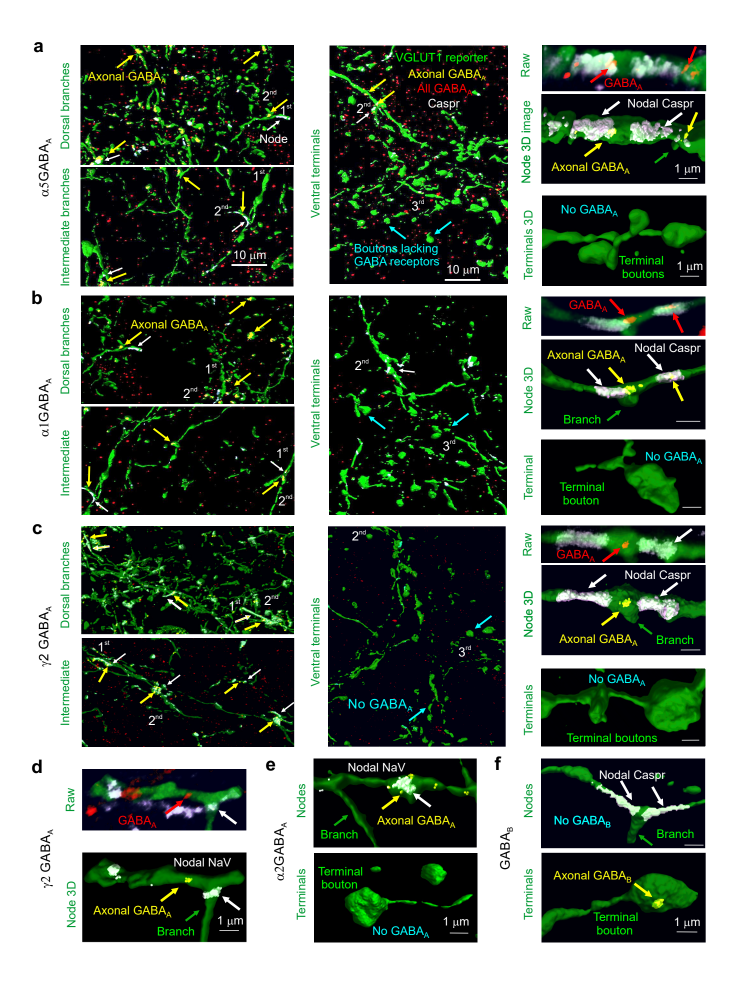
Fig. 7 | Facilitation of reflexes in humans. a-b, MSR in soleus EMG evoked by a tibial nerve pulse (1.1xT, 0.2 Hz, bi), and phasic facilitation of the MSR following a brief conditioning of the cutaneous branch of the deep peroneal nerve (cDP nerve) at varying intervals (ISIs, bii, 1.0xT, perception threshold T, at rest), and lack of changes in background (Bkg) motor unit (MU) activity or EMG evoked by conditioning alone (biii, peri-stimulus frequencygram, PSF; with weak contraction). c, Same as (b), but with proprioceptive conditioning evoked by a brief tibial anterior (TA) muscle tendon vibration, which alone inhibited MU activity (postsynaptic inhibition, PSF Bkg, ciii). d, Summary box plots of changes in MSR and postsynaptic (MU) activity with brief conditioning (cDP, n = 14 subjects; or TA vibration, n = 6; as in **b-c**) and long cutaneous conditioning trains (e, n = 14). * significant change with conditioning, P < 0.05. e, Tonic increase in MSR (tonic facilitation) after 0.5 s cutaneous conditioning train (cDP, 1.1xT, 200 Hz) at rest (ei-ii), without prolonged changes in MU activity induced by conditioning alone (eiii, PSF in weak contraction). MSR evoked by tibial stimulation every 5 s, with averages from repeated conditioning shown in (eii). * significant change in MSR, P < 0.05, n = 5subjects. f, Overlay of all MU firing rates (PSF) with repeated MSR testing (at 5 s intervals) during ongoing weak contraction, and effect of the 0.5 s cutaneous conditioning train (fi). Summary box plots of increased probability of MU firing during MSR (fii), without changing estimated EPSP size (fiii, PSF thin line; thick line unitary EPSP shape from Fig. 5i) or background MU firing (Bkg, fiv). * significant change with conditioning, P < 0.05, n = 10 subjects.

Supplementary information

GABA facilitates spike propagation at branch points of sensory axons in the spinal cord

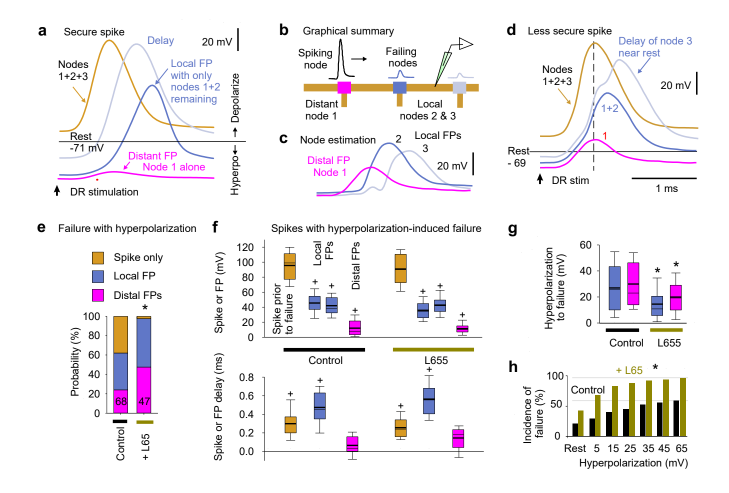
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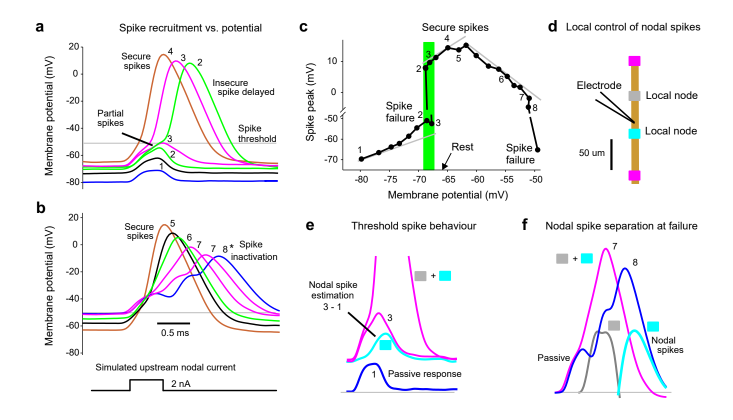
Extended Data Fig. 1 | Nodal and not terminal GABA_A receptors in mice.

a-c, In the sacrocaudal spinal cord of mice we examined the distribution of GABAA receptor subunits on nodes and terminals of sensory axons, including extrasynaptic α 5 subunits, synaptic α 1 and α 2 subunits, and ubiquitous γ 2 subunits (e.g. forming the common $\alpha 1\beta \gamma 2$ or $\alpha 5\beta \gamma 2$ receptors, though less common extrasynaptic $\alpha 1\beta \delta$ have been reported)^{32,108}-¹¹⁰. We genetically labelled primary sensory axons by their expression of the vesicular glutamate transporter VGLUT1 with a reporter in VGLUT1^{Cre/+}; R26^{LSS-tdTom} mice (tdTom reporter displayed as green, for consistency with Fig. 1). VGLUT1 is mainly only expressed in sensory axons⁵⁹, especially ventral proprioceptive afferents, as other afferents do not reach the ventral horn¹⁶. Axons were reconstructed in 3D and branching order is indicated (1st, 2nd and 3rd), as detailed in Fig. 1. Typical axon branching patterns are shown in the deep dorsal horn (left), intermedial laminae and ventral horn (middle). Expanded images of nodes of Ranvier identified by paranodal Capsr immunolabelling are shown on 1st order branches in the intermediate laminae (on right), along with raw confocal images prior to 3D reconstruction. Expanded terminal boutons from the ventral horn are also shown on the right. As in rats, nodes are often near branch points (where 2nd order branch arises from 1st order branches in this case, green arrow). We identified axonal GABA receptors as those colocalized with axonal tdTom (yellow, as in Fig. 1), and the remaining GABA receptors are labelled red in the 3D reconstructions. The α 5, α1, and γ2 GABA_A receptors subunits were found on large axon branches (1st and 2nd order) in the dorsal, intermediate, and ventral cord, near nodes counterstained with Caspr immunolabelling (white, arrows), but not on ventral horn terminal boutons (3rd order; examples at blue arrows; expanded on right). The presence of these subunits is consistent with their mRNA previously reported in the dorsal root ganglion¹¹¹. Also, the finding of $\alpha 1$ subunits on these axons is consistent with the recent observation that $\alpha 1$ is only on myelinated sensory axons, rather than unmyelinated C fibres¹¹². Similar results were obtained from n = 3 mice. All terminals are VGLUT1⁺ (not shown). **d**, Immunolabelling for $\gamma 2$ GABA_A receptor subunits on 3D reconstructed sensory axons, with same format and mice of (a), but with nodes identified by Nav (sodium channels), rather than Caspr. Again, the node and GABA receptors are near a branch point (green arrow). e, Immunolabelling for α 2 GABA_A receptors on 3D reconstructed sensory axons, with same format and mice of (a), but nodes identified by Na_V. Again, GABA_A receptors were near nodes and not on terminals. f, Immunolabelling for GABA_B receptors on 3D reconstructed sensory axons, with same format and mice of (a). GABA_B receptors were generally absent from nodes identified by Caspr, but present on ventral terminal boutons, as in rats (Fig. 1).



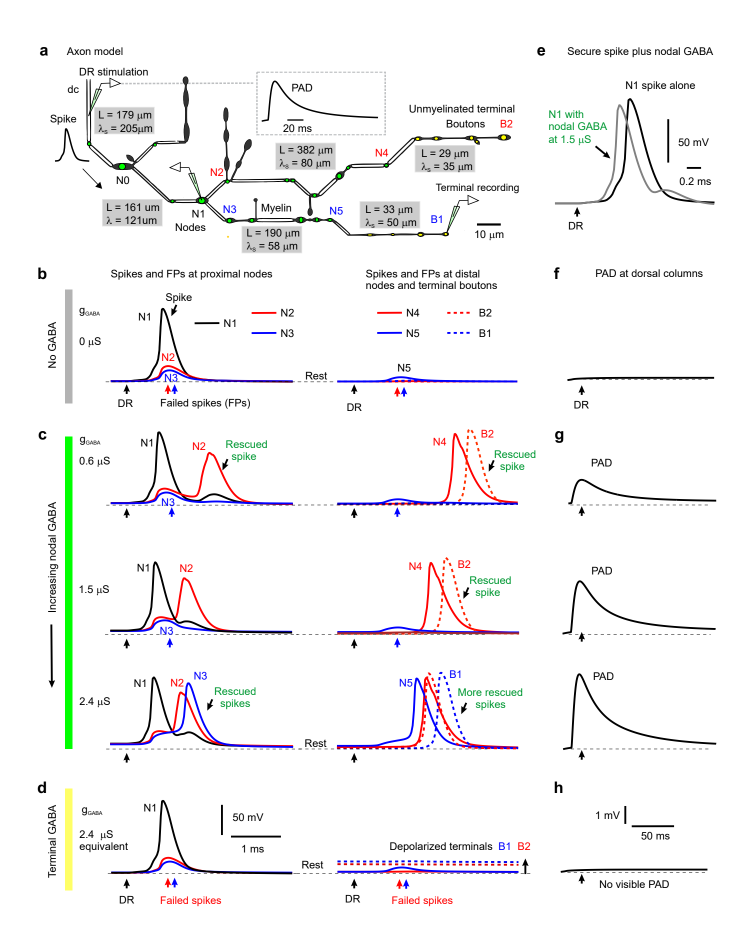
Extended Data Fig. 2 | Voltage dependence of nodal spike propagation failure following dorsal root stimulation.

a-d, Intracellular recording from proprioceptive la afferent branches in the dorsal horn with secure spikes at rest, evoked by DR stimulation (1.1xT, 0.1 ms; sacral S4 DR; a, d). Spike failure was induced by increasing hyperpolarization (failure near rest in d, but not a), with a delay and then abrupt loss of height, reflecting failure of successively further away nodes (a, d). Of the two local nodes adjacent to the electrode, one failed first with hyperpolarization, leaving the attenuated spike from the other node (local FP, about 1 λ_s away), which eventually failed as well with further hyperpolarization, leaving a much smaller FP from more distal nodes (distal FP). Spike failure in a node was always proceeded by a delay in the nodal spike. Estimated contributions from local nodes (b-c) were computed by subtraction from traces in (d). Note that due to attenuation of injected current with distance, larger hyperpolarizations were needed to stop spikes with more distal vulnerable nodes (secure spikes), as observed by smaller distal FPs (a verses d), and some axon spikes could not be stopped (not shown; Spike only, quantified in e). Data collected in discontinuous current clamp (DCC) mode so electrode rectification during the injected hyperpolarizing current did not affect potential (DCC switching rate 7 kHz, low pass filtered at 3 kHz to remove switching artifact, which also removed stimulus artifact). e, Distribution of spiking in branches with full spikes only, one local nodal spike (local FP), or a distal nodal spike (distant FP) remaining after maximal hyperpolarization, and * significant change when blocking $\alpha 5$ GABA_A receptors with L655708 (0.1 - 0.3 μ M) using γ -squared test, P < 0.05; n =68 control and n = 47 L655708 treated branches. f, Box plots of spike or FP heights and delays in branches indicated in (e), measured just prior to spike failure (or at maximal hyperpolarization for secure spikes) and after failure (for local and distal FPs, as induced in a-d). Delay measured relative to peak of spike at rest. + FP significantly different than spike at rest, P < 0.05. g, Box plots of the hyperpolarization needed to induce failure, in branches from (e). * significant change with L655708, P < 0.05. h, Incidence of failure at varying potentials (% of total spikes from e). * significant change with L655708, χ -squared test, P < 0.05.

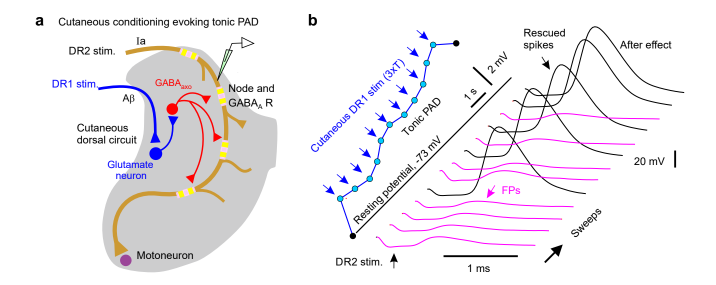


Extended Data Fig. 3 | Voltage dependence of spike failure with simulated nodal currents.

Simulating spike propagation failure in a proprioceptive axon by applying a brief intracellular current injection to mimic the current arriving from an upstream node (and FP), yielded full spikes evoked at rest, but nearby nodal spikes delayed and then failing as the membrane was held progressively more hyperpolarized with a steady bias current. Also, large steady depolarizations inactivated these spikes, though well outside of the physiological range (> -50 mV). a, Intracellular recording from proprioceptive Ia afferent branch in the dorsal horn (sacral S4 axon). A brief current injection pulse (0.5 ms) was applied to simulate the current arriving from distal nodes during normal axon spike conduction, and repeated at 1s intervals. During these pulses the membrane was held at varying potentials for 1-2 s with steady current injection, with numbers and colours denoting a given holding potential (in DCC mode). At the most hyperpolarized levels spikes failed to be evoked and only the passive response is seen, like a FP (blue, 1). As the potential was depolarized to near the axon's resting potential (-67 mV) partial spikes occurred (green, 2 and 3), likely from a single adjacent node activating, and then delayed broad spikes occurred, as both adjacent nodes were activated. At more depolarized levels the spikes arose more rapidly and increased in height to full secure spikes (4). b, In the same axon as (a), at holding potentials well above those seen physiologically (near -50 mV, lower plots) spikes started to exhibit sodium channel inactivation and failure, with a decrease in spike height and delay (7 - 8) and eventually full failure (shown in c). Adjacent nodes started failing at slightly different times with different delays, broadening the spike and eventually separating into two distinct nodal spikes (8*). c, Spike heights plotted as a function of holding potential, including those spikes illustrated in (a and b), with spike numberlabels indicated. Left grey line indicates passive leak current response, and shows deviation from passive response near rest. Shaded green region shows all or nothing failure or spikes near the resting potential. Middle grey line shows a region of secure spikes with relatively invariant spikes. Right grey line shows spike inactivation with large depolarizations and outright failure near - 50 mV. Note the split vertical axis. Similar voltage dependence of spike failure occurred for n = 5/5axons tested. This demonstrates two modes of spike failure: 1) spikes that fail at rest or at hyperpolarized potentials and 2) spikes that fail with large depolarizations above rest. The latter is likely not physiological, since even the largest PAD that we have observed (5 - 10 mV; Fig. 4d) does not depolarize axons to -50 mV, since axons rest near -70 mV, and PAD is only large at hyperpolarized levels (Fig. 4d) and decreases steeply as the potential approaches the reversal potential for chloride (-15 mV)¹⁶. d, Schematic of recording arrangement and relation to adjacent nodes for data in (e) and (f). e, Expansion of responses 1 (blue) and 2 (pink) from (a), and difference (cyan) to show the first local nodal spike height at threshold, recorded at electrode. Active nodes from schematic in (d) shown with shaded boxes. f, Spikes near sodium inactivation from (b) (7 and 8), with differences indicating local nodal spikes (cyan and grey, nodes from d).

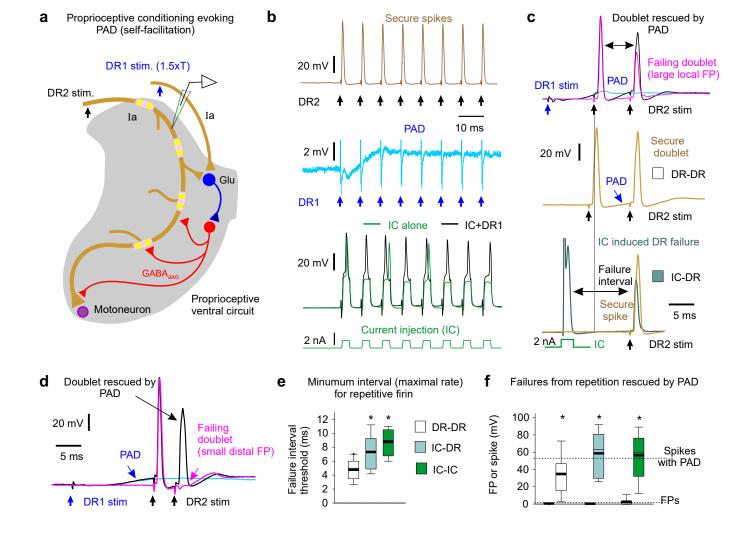


Extended Data Fig. 4 | Computer simulation of branch point failure and rescue by GABA. a, Model of a 3D reconstructed proprioceptive afferent, drawn to scale, except myelinated branch lengths all shortened an order of magnitude. Double line segments are myelinated (white) and the rest unmyelinated. Adapted from anatomical studies of Walmsley³⁴. Nodes are indicated with a green dot, and ventrally projecting terminal boutons indicated with a yellow dot. As in our axons of Fig. 1, branch points were always at nodes. GABAA receptors of equal conductance (nS) were placed at each branch point (node), and total dorsal columns (dc) depolarization from phasically activating these receptors is shown in inset. The branch lengths (L) and computed space constants (λ_s) are indicated in gray boxes for each segment of the afferent, the latter computed from subthreshold current injections into each segment. From left to right the gray boxes are for segments spanning from the dorsal columns (dc) to N0, N0 to N1, N1 to N5, N1 to N4, N4 to B2 and N5 to B1. Average space constant was $\lambda_S = 91 \, \mu m$, similar to in other axons³⁹. **b**, Responses to simulated dorsal root (DR) stimulation (0.1 ms pulse, 2 nA, at black arrows) computed at various downstream branch points (nodes) and terminal boutons in the spinal cord, with resting potential indicated by thin dashed line (-82 mV). A sodium spike propagated to the branch point at node N1, but failed to invade into the downstream branches, leaving nodes N2 and N3 with only a passive depolarization from the N1 spike (failure potential, FP). Further downstream nodes and terminals experienced vary little depolarization during this failure (N5, B1 and B2). In this case the GABA conductance was set to zero (g_{GABA} = 0, control), simulating a lack of GABA tone. Note that only the nodes beyond the parent branch at node N1 failed due to the conductance increases in large daughter branches to nodes N2 and N3 that drew more current than node N1 could provide (shunting conductances). Node 3 is particularly complex with a large unmyelinated region (black) and a nearby branch contributing to its conductance and related failure, and Node 2 has two adjacent branches contributing to its conductance. Other branch points with relatively smaller conductance increases (NO, simpler branching) did not fail to conduct spikes. Generally speaking, if the upstream node of a parent branch cannot provide enough current to activate the nodes of its daughter branches then spikes fail, and this is especially likely with multiple sequential branch points, like in N1 – N2. \mathbf{c} , As seen experimentally, nodal GABA_A receptor activation to produce PAD prior to the DR stimulation (~ 10 ms prior; as in Fig. 4f) rescued spikes from failing to propagate. That is, with GABA_A receptors placed just at branch points (i.e., at nodes) a weak phasic activation of these receptors (conductance $g_{GABA} = 0.6 - 1.5 \,\mu\text{S}$ per node shown) rescued conduction down the branch to node N2, with full nodal spikes seen at the distal node N4 and the terminal bouton B2 (DR stimulation at peak of PAD, detailed in (g) with black arrows indicating DR stimulation timing). A larger GABA receptor activation (2.4 nS) additionally rescued spike conduction down the branch to node N3, with full spike conduction to the distal node N5 and the terminal bouton B1. Note that increasing GABA conductance sped up the arrival of distal spikes (e.g. at N4 and B2), by up to 1 ms, suggesting substantial variation in sensory transmission times induced by GABA, as we see experimentally. Also note that this nodal GABA depolarized the nodes (N1 - N3) relative to rest (thin dashed line), thus assisting spike initiation. In contrast, nodal GABA did not depolarize the terminal boutons (B1 and B2), consistent with our recent direct recordings from terminals¹⁶. Sensitivity analysis revealed similar results with a wide range of sodium channel and GABA receptor conductances, though increasing sodium conductance sufficiently prevented failure all together (not shown, but like in Extended Data Fig. 9f). d, When we removed all nodal GABA receptors and instead place them on terminal boutons (near B1 and B2, yellow, with equivalent total conductance, 2.4 µS condition), then activating them did not rescue the spike propagation failure, since the associated depolarization of nodes is too attenuated at the failure point (N1-N3; no change from resting potential). The GABA receptors did depolarize the terminal boutons (B1 and B2, thick dashed lines) substantially relative to the resting potential (thin dashed lines), but this depolarization was sharply attenuated in more proximal nodes (N1-3). e, Reduction of spike height (shunt) and speeding of spike onset with increasing GABA conductance at a non-failing node (N1; model with nodal and not terminal bouton GABA conductances, c), consistent with actual recordings from axons in Fig. 3d and Extended Data Fig. 7a. f-h, PAD recorded at the dorsal columns (dc) during conditions in (b-d), respectively, as experimentally recorded dorsal PAD. A phasic GABA induced depolarization (PAD) was generated by changing GABA conductances, g_{GABA}, as detailed in Methods, and GABA receptor location varied as in (c-d). DRs were stimulated at the peak of this PAD in (c-d). Note that nodal (g) but not terminal (h) GABA_A receptors caused a visible depolarization (PAD) at the dc, due to less electrotonic attenuation over a shorter distance to the dc.



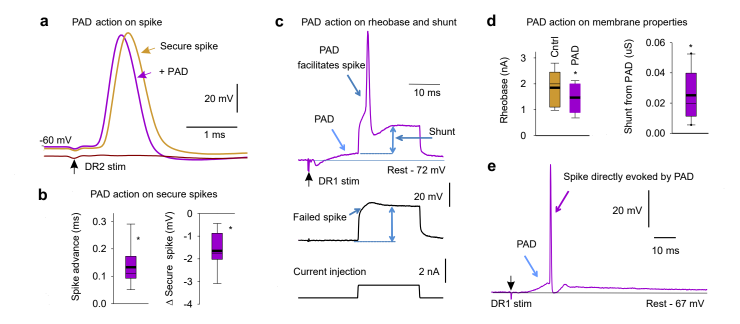
Extended Data Fig. 5 | Cutaneous driven trisynaptic circuits mediating PAD and assisting repetitive firing.

a, Cutaneous driven dorsal trisynaptic circuit mediating PAD. A minimally trisynaptic circuit is classically known to depolarize afferents via GABA_{axo} neurons. This circuit involves sensory afferents activating excitatory intermediary neurons (glutamatergic) that in turn activate GABA_{axo} neurons that return to innervate sensory axons¹⁴. Even though GABA_{axo} neurons are small⁶ this circuit influences afferents over widespread regions of the spinal cord¹⁶. Specifically, the activation of a small group of sensory axons in just one DR or nerve causes this trisynaptic circuit to produce a widespread activation of many axons across the spinal cord, even many segments away and across the midline¹⁶. This allowed us to activate PAD from adjacent roots without directly activating an axon in a particular root, as detailed in Fig. 4 and the rest of this figure. One variant of this classic trisynaptic circuit specifically involves cutaneous stimulation activating dorsal intermediary neurons^{14,15} that activates GABA_{axo} neurons that in turn innervate cutaneous afferents, which we term the cutaneous dorsal circuit. While this cutaneous dorsal circuit also synaptically innervates some proprioceptive afferents¹¹³ (a), its main action on proprioceptive afferents is to produce a pronounced extrasynaptic spillover of GABA that depolarizes these afferents tonically via α5 GABA_A receptors (termed: tonic PAD, L655708 sensitive), especially with repetitive cutaneous nerve stimulation (1 - 200 Hz) that leads to minutes of depolarization¹⁶, and we see similar tonic PAD here (detailed next). b, Intracellular recording from a proprioceptive axon branch in rat dorsal horn (sacral S4 axon, DR2). The axon branch spontaneously exhibited spike propagation failure when its was stimulated alone (denoted DR2 stimulation, repeated at 1 Hz, 1.1xT, 0.1 ms), with only a small failure potential (FP) visible (lower pink traces). Activation of a largely cutaneous DR (caudal Ca1 DR, innervating the tip of the tail, stimulation at intensity for cutaneous afferents, 3xT, 0.1 ms; denoted DR1) evoked a slowly rising tonic PAD when repeated at 1 Hz (blue). When the axon stimulation (DR2 stimulation) was combined with the repeated cutaneous stimulation (DR1, 60 ms prior to each DR2 stimulation) the slowly building PAD prevented spike failure (black spikes), and this outlasted the cutaneous stimulation (after effect). Similar results obtained in n = 20/20axons tested.



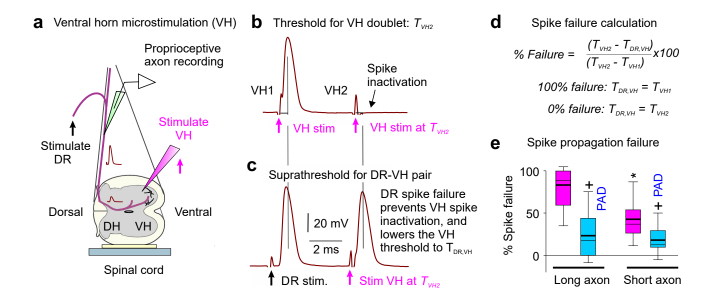
Extended Data Fig. 6 | Proprioceptive driven trisynaptic circuit for PAD enabling high frequency spike transmission.

a, Proprioceptive driven ventral trisynaptic circuit mediating PAD. Another variant of the classic trisynaptic circuit involves proprioceptive afferents activating excitatory intermediary neurons (glutamatergic) that then activate GABA_{axo} neurons that innervate these same afferents, including ventral terminal regions of the afferents^{14,16}. This circuit is more ventrally located compared to the cutaneous dorsal PAD circuit¹⁴, and thus, we term it the proprioceptive ventral PAD circuit. Importantly, the circuit is recurrent, with proprioceptive afferents causing self-facilitation of themselves (via homonymous PAD). It produces a fast phasic axon depolarization (phasic PAD, fast synaptic; Fig. 4c), as well as a slower tonic depolarization (tonic PAD, Fig. 4c, likely from extrasynaptic GABA spillover), as detailed previously¹⁶. Since proprioceptive sensory axons naturally fire at high rates¹¹⁴ where they are vulnerable to spike failure (Fig. 2e-f), we examined the action of self-facilitation by GABA on this failure. b-f, During rapid repetitive stimulation of a DR to evoke spikes in a proprioceptive axon there was an inevitable activation of PAD from low threshold proprioceptive axons (homonymous PAD, b). This PAD helped spikes fire at high physiological rates of up to 200-300 Hz (5 – 3 ms intervals) before spike inactivation and failure occurred because, in absence of PAD, isolated repetitive activation of the axon with intracellular current pulses (IC) led to failure at much lower rates (~100 Hz; longer spike intervals; b, e), even after just two stimuli (doublets; c, e). Additional PAD evoked by simultaneous stimulation of an adjacent DR (2xT) reduced failure from fast repeated IC stimuli (b, f), repeated DR stimuli (doublet, c, d, f) or hybrid IC-DR stimulation pairs (f). Legend details are as follows: b, Intracellular recording from proprioceptive axon in rat dorsal horn (sacral S4 axon) with spikes securely evoked by fast repeated DR stimulation (top, 1.1xT, 0.1 ms, sacral S4 DR, denoted DR2; resting potential -68 mV), but spikes failing intermittently with repeated intracellular current injection (IC) at the same rate (bottom green), due to sodium channel inactivation. The reason that spike failure does not occur with the fast DR stimulation is that it is accompanied by a build up of tonic PAD (from self-activation) that helps prevent failure, because adding to the IC stimulation a simultaneous conditioning stimulation of other proprioceptive afferents in an adjacent DR (DR1 stim at 1.5xT, 0.1 ms, S3 DR) prevents spike failure (black trace, bottom), via the proprioceptive ventral circuit (a). This DR1 conditioning stimulation does not directly activate spikes in the axon, but it causes a fast depolarization (phasic PAD) that rapidly helps spikes (as early as 6 - 10 ms later), and a building tonic depolarization (tonic PAD) with repetition that further helps later spikes in the stimulation train (DR1 stimulation alone blue, middle trace). Similar results obtained in n = 7/7 axons. Likely similar tonic PAD and associated increased spike conduction helps explain post-tetanic potentiation of the monosynaptic EPSP, as previously suggested²⁹. c, Repeated DR stimulation at higher rates eventually causes spike failure in proprioceptive axons (sodium spike relatively refractory), as shown in the top panel where a double stimulation (doublet, S4 DR, denoted DR2-DR2, 1.1xT, 0.1 ms, resting at -75 mV) exhibits failure on the second spike (with large FP indicated, magenta). However, additional PAD provided by stimulating an adjacent DR (DR1; 1.5xT, group I intensity, 0.1 ms) about 10 ms earlier helps prevent this spike failure (black trace; blue trace: PAD alone). When the same axon was stimulated slightly slower (with a longer doublet interval, second plot, DR-DR) failure did not occur, which we designate the failure interval threshold, which is quantified in (e). The selfactivated PAD caused by the first DR stimulation in this doublet helped prevent failure in second DR stimulation because replacing the first DR stimulation with an intracellular stimulation (IC, 2 nA) to activate the spike leads to failure of the second spike evoked by the DR stimulation at much longer intervals (lower trace, IC-DR). d, Another example of a failed doublet spike (DR2-DR2 stim, 1.1xT, 0.1 ms) that was rescued by PAD evoked by adjacent DR stimulation (DR1 1.5xT, 0.1 ms, resting at -78 mV), as in the top plots of (c), except that in this case the failure is at a more distal node, since the FP is small. e, Failure interval threshold (minimum firing interval prior to failure, or maximal firing rate) with DR doublets (DR-DR), IC doublets (IC-IC) or IC-DR pair stimulation. Note the shorter intervals possible with the PAD evoked by the first stimulation (DR-DR). * significantly longer than minimum DR doublet interval (DR-DR), n = 18 each, P < 0.05. f, Quantification of the FP heights that were induced by a fast doublet (DR-DR or IC-IC; n = 14 each, at failure threshold interval) or IC and DR stimulation (IC-DR; n = 10), and the rescue of spikes by PAD evoked by adjacent DR stimulation. * significant increase in height with PAD, P < 0.05.



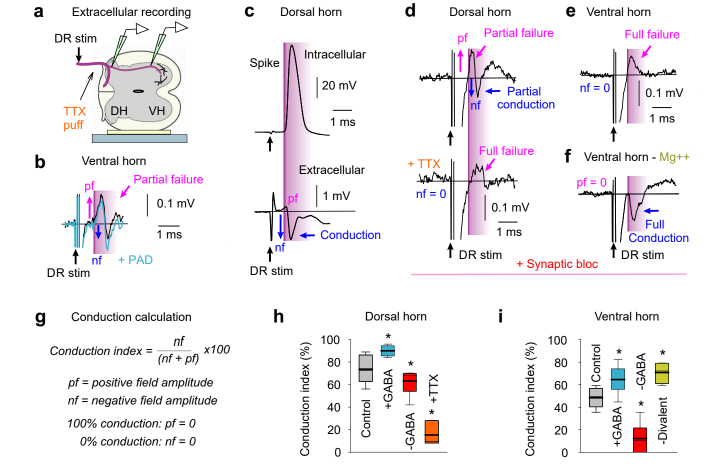
Extended Data Fig. 7 | Other excitatory actions of GABA_A receptors on proprioceptive axons.

a, Intracellular recording from a sacral S3 proprioceptive la afferent branch in the dorsal horn with a secure spike evoked by S3 DR stimulation at rest (DR2, 1.1xT, 0.1 ms, - 60 mV rest, rat). Sensory-evoked PAD initiated by stimulating an adjacent DR (DR1; S4 DR; 2xT, 0.1 ms pulse, as in Fig. 4) 10 ms prior to the DR2 stimulation only moderately influenced the spike (DR1 stimulation time not shown). It sped up the spike latency and rise time, reduced the fall time and slightly reduced the spike height. Hyperpolarization induced spike failure (lower trace), as in Extend Data Fig. 2a. b, Summary box plots of change in spike peak latency (advance) and height with prior sensory PAD activation as in (a). *significant change, P < 0.05, n = 20 and 26, respectively. c, Intracellular recording from an S3 proprioceptive Ia afferent branch in the dorsal horn with a brief current injection pulse just subthreshold to initiating a spike (near rheobase, bottom trace) at rest, only initiating a passive response with a small failed spike (middle trace). However, prior activation of PAD by stimulating an adjacent DR (DR1 as in A; S4 DR, 2xT, 0.1ms top trace) allowed the same current pulse to evoke a spike (above rheobase). The passive response to the current injection (double blue arrow; resistance R = V/I) was decreased during the PAD, corresponding to an increase in conductance, that contributed to a shunt (reduction) of the currents generating the spike, though this only caused about a 1% drop in spike height (1 mV; b). DCC recording mode, as in Extend Data Fig. 2a-d. d, Summary box plots of rheobase (current threshold from c) before and during PAD, and change in shunt (conductance = 1/R) with PAD, as in (c). * significant change with PAD, P < 0.05, n = 37 axon branches. e, By itself sensory evoked PAD sometimes initiated a spike on its rising phase, when the DR stimulation was large enough, demonstrating a direct excitatory action of GABAA receptors, as previously reported¹⁶. These spikes propagate antidromically toward the DR; and are thus termed dorsal root reflexes (DRR). Example intracellular recording from rat sacral S3 proprioceptive afferent branch in the dorsal horn, similar to in Fig. 4e (top trace), but PAD evoked by a larger DR1 stimulation (S4 DR stimulation 3.5xT, 0.1 ms) that itself evoked a spike (DRR). These PAD evoked spikes occur with a variable latency of $10 - 30 \text{ ms}^{16}$ and thus make axons refractory for about 30 ms after the DR stimulation¹¹⁵. These spikes can also evoke EPSPs in motoneurons via the monosynaptic pathway⁹, and thus also produce a post activation depression of the EPSPs for many seconds. We thus kept the PAD low when examining the effects of PAD on EPSPs, to avoid these spikes and their subsequent inhibitory action (in Figs. 5-7).



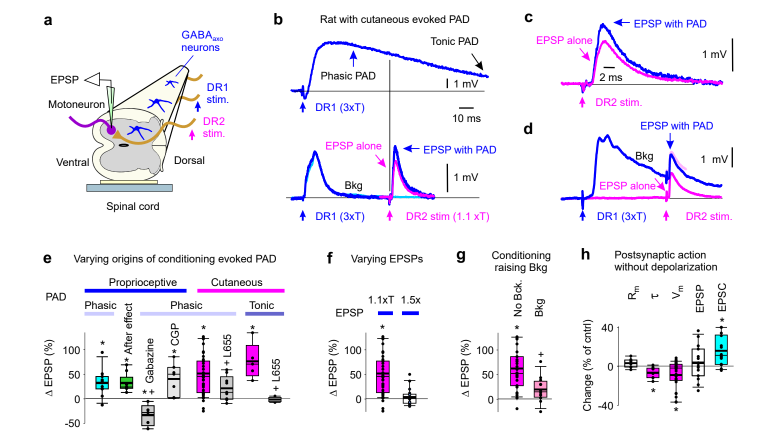
Extended Data Fig. 8 | Estimating the overall spike conduction failure from the dorsal root to the motoneurons.

a, Experimental setup to indirectly measure sensory axon conduction failure following DR stimulation, by examining whether failed axon segments are relatively less refractory to activation after failure, using a double pulse method adapted from Wall³⁰. A tungsten microelectrode (12 $M\Omega$) was placed in the ventral horn (VH) near the sensory axon terminals on motoneurons (S3 or S4 VH), to activate the branches/nodes of the axon projecting to the motoneuron that may have failed (VH stimulation). Spikes from VH or DR stimulation were recorded intracellularly in a proprioceptive axon penetrated in the dorsal columns. b, VH threshold in refractory period. Rapidly repeated VH stimulation (VH doublet; two 0.1 ms pulses) at an interval short enough to produce spike inactivation on the second stimulation (4 ms), with stimulus current adjusted to threshold for inactivation, T_{VH2}. This T_{VH2} (~15 uA) was always higher than the threshold VH stimulation for evoking a spike with the first stimulation, T_{VH1} (~10 uA, not shown). Recorded in sacral S4 afferent resting at -72 mV, with doublets repeated at 3 s intervals to determine current thresholds. c, VH threshold after DR stimulation. Similar repeated activation of the axon in (b), but with the first activation from a DR stimulation (at 1.5x DR threshold) and the second from VH stimulation at the T_{VH2} intensity from (b). In this case the VH stimulation readily activated the axon spike, likely because the DR-evoked spike did not propagate to the VH, leaving the silent portion of the axon non refractory. Thus, this VH stimulation evoked spikes with a lower current than T_{VH2} , with this lower threshold denoted $T_{DR,VH}$ ($\sim 12 \mu A$, not shown). This DR – VH stimulation interval was deliberately set too short for the involvement of PAD (which rises in > 4 ms; Fig. 4). d, Computation of spike failure based on changes in VH stimulation thresholds. If the DR-evoked spike entirely fails to propagate to the VH, then the threshold for subsequently activating the VH (TDR,VH) should be the same as the threshold without any prior activation (T_{VH1} = T_{DR,VH}), whereas if it does not fail then the threshold for activating the VH should be the same as with a VH doublet ($T_{VH2} = T_{DR,VH}$). In between these two extreme scenarios, the DR evoked spike may only partially fail to propagate spikes to the VH; in this case $T_{DR,VH}$ should be between T_{VH1} and T_{VH2} , with the difference T_{VH2} - T_{VH1} representing the range of possible thresholds between full failure and conduction. Overall the % conduction failure can be thus quantified as: $(T_{VH2} - T_{DR,VH})/(T_{VH2} - T_{VH1}) * 100\%$, which is 100% at full failure and 0% with no failure. e, Average spike conduction failure to the VH in proprioceptive axons, and decrease following a DR conditioning stimulation that depolarized the axon (PAD). Box plots of failure estimated as in (b-d). Prior DR conditioning to produce PAD (via adjacent S4 or Ca1 DR stimulation at 3xT) reduced the failure estimated 20 ms later by the paired-pulse conduction testing (repeating DR – VH stimulations of b-d). DR conditioning itself lowered the thresholds for VH activation, as previously reported (not shown)³¹. We studied two lengths of axons: long axons (intersegmental, n = 10) with the VH stimulation one segment away from the recording site, and short axons (segmental, n = 12) with the VH stimulation near the recording site, in the same segment. + significantly less failure with PAD and * significantly less failure with short compared to long axons, *P* < 0.05.



Extended Data Fig. 9 | Axonal spike conduction estimated from extracellular recordings.

a, Experimental setup to directly record spike conduction failure in proprioceptive axon terminals in the ventral horn (VH) following DR stimulation. Extracellular (EC) recordings from axon terminals in VH, with glass electrode positioned just outside these axons, and for comparison EC recording in the dorsal horn (DH). b, EC field recorded in VH after DR stimulation (S4 DR, 1.1xT), with a relatively large initial positive field (magenta arrows, pf) resulting from passively conducted axial current from sodium spikes at distant nodes (closer to the DR; outward current at electrode), some of which fail to propagate spikes to the VH recording site; thus, this field is a measure of conduction failure, as demonstrated in (c-f) below. Following this, a negative field arises (blue arrow, nf), resulting from spikes arising at nodes near the electrode (inward current); thus, this field is a measure of secure conduction. Reducing conduction failure by depolarizing the axon (+PAD) with a prior conditioning stimulation of an adjacent DR (Ca1, 2xT, 30 ms prior), decreased the positive field (pf) and increased the negative field (nf), consistent with increased conduction to the terminals, and in retrospect the same as Sypert et al. (1980) saw in cat (their Fig. 4). Large stimulus artifacts prior to these fields are truncated. c, Control recordings from proprioceptive axons in dorsal horn (DH) to confirm the relation of the EC negative field (nf) to spike conduction. Intracellular (IC) recording from axon (sacral S4, resting at -64 mV) and EC recording just outside the same axon, showing the DR evoked spike (IC) arriving at about the time of the negative EC field (nf). There is likely little spike failure in this axon or nearby axons, due to the very small initial positive field (pf). EC fields are larger in DH compared to VH (G, 10x), and thus the artifact is relatively smaller. d, Locally blocking nodes with TTX to confirm the relation of the positive EC field to spike failure. EC recording from proprioceptive axon in the dorsal horn (S4), with an initial positive field (pf) followed by a negative field (nf), indicative of mixed failure and conduction. A local puff of TTX (10 μl of 100 μM) on the DR just adjacent to the recording site to transiently block DR conduction eliminated the negative field (nf) and broadened the positive field (pf), consistent with distal nodes upstream of the TTX block generating the positive field via passive axial current conduction, and closer nodes not spiking. Recordings were in the presence of synaptic blockade (with glutamate receptor blockers, kynurenic acid, CNQX and APV, at doses of 1000, 100 and 50 μM respectively), to prevent TTX spillover having an indirect action by blocking neuronal circuit activity, including GABAaxo neuron activity. This synaptic blockade itself contributed to some spike failure, consistent with a block of GABA_{axo} neuron activity, as there was a more prominent positive field (pf) compared to without blockade in (c). e, EC field recorded from terminals of proprioceptive axons in the ventral horn near motoneurons (S4), in the presence of an excitatory synaptic block that largely eliminates most neuronal circuit behaviour (with kynurenic acid, CNQX and APV, as in d). In this synaptic block negative fields were generally absent (nf = 0), and only prominent positive fields (pf) occurred (as with TTX block), suggesting that conduction to the VH often completely failed when circuit behavior is blocked, which likely indirectly reduces GABA_{axo} neuronal circuit activity and its associated facilitation of nodal conduction. f, Rescue of spike conduction to the ventral horn by increasing sodium channel excitability by reducing the divalent cations Mg++ and Ca ++ in the bath medium⁷⁵. Same EC field recording as in e, but with divalent cations reduced (Mg++, 0 mM; Ca++, 0.1 mM). The positive field was largely eliminated (pf = 0) and replaced by a negative field (nf), consistent with elimination of conduction failure, and proving that the positive field is not a trivial property of axon terminals 73,74,116. g, Conduction index computed from positive (pf) and negative (nf) field amplitudes as: nf / (nf + pf) x 100%, which approaches 100% for full conduction (pf \sim 0; as in c) and 0% for no conduction (nf = 0; as in e). h-i, Summary of conduction index estimated from EC field potentials, shown with box plots. Without drugs present in the recording chamber, the axon conduction from the DR to the dorsal horn is about is 70% (h, n = 17), consistent with Fig. 2, whereas conduction from the DR to the VH is only about 50% (i, n = 9), suggesting substantial failure at the many branch points in the axon projections from the dorsal horn to the motoneurons. Increasing GABAaxo neuron activity with DR conditioning (30 – 60 ms prior) increased conduction (+GABA, in both the DH and VH, n = 5 and 9, as in b), whereas decreasing GABA and all circuit activity in a synaptic blockade decreased conduction (-GABA, in both DH and VH, n = 5 and 6, as in **d-e**). TTX (n = 4, **h**) or removal of divalent cations (Mg⁺⁺ and Ca⁺⁺, -Divalent, n = 4, **i**) reduced or increased conduction, respectively (as in **d** and **f**). * significant difference from control pre-drug or pre-conditioning, P < 0.05.



Extended Data Fig. 10 | Sensory evoked facilitation of monosynaptic EPSPs by GABA_A receptors.

a, Whole spinal cord ex vivo preparation for intracellular recording of EPSPs from motoneurons while stimulating dorsal roots (DRs). b, Monosynaptic EPSP in an S4 motoneuron evoked by a proprioceptive group I stimulation of the S4 DR (1.1xT, denoted DR2, lower traces; resting potential -75 mV: black line), alone (pink) and 60 ms after (blue) a conditioning stimulation of cutaneous afferents in rat to evoked PAD (stimulation of the largely cutaneous Ca1 DR, 2.5xT; denoted DR1). Averages of 10 trials each at 10 s intervals. PAD evoked by the same cutaneous conditioning stimulation in a proprioceptive S4 DR afferent is shown for reference (top, recorded separately, as in Fig. 4b). c, EPSPs from (b) on expanded time scale. d, Similar to (b), but stronger conditioning stimulation (DR1, 3xT) evoking background postsynaptic activity (blue, Bkg) that lasted longer than 60 ms, and slightly inhibited the EPSP, likely from increased postsynaptic conductances shunting the EPSP (postsynaptic inhibition; light pink: overlay of EPSP alone) and masking nodal facilitation. e, Summary box plots of facilitation of EPSPs during phasic PAD evoked by either proprioceptive conditioning (S3 or contralateral S4 DR stimulation, 1.1xT, in rats and mice similar and combined, n = 6 and 5, blue) or cutaneous conditioning (Ca1 DR stimulation, 2-3xT, in rats, n = 42, pink), and action of GABA_A and GABA_B antagonists (gabazine 50 μ M, L655708 0.3 μ M and CGP55845 0.3 μ M, grey, n = 5 - 9 each). EPSPs evoked in S3 and S4 motoneurons by DR2 (S3 or S4) stimulation at 1.1T, as in (b). Facilitation measured 60 ms post conditioning during phasic PAD (phasic condition indicated) and when postsynaptic actions of conditioning (Bkg) were minimal (as in b). After conditioning was completed EPSP testing continued and revealed a residual facilitation that lasted for 10 - 100 s (After effect, green, n = 9 tested), due to a build up of tonic PAD, after which the EPSP returned to baseline (not shown), similar to post-tetanic potentiation²⁹. Also, a brief high frequency cutaneous stimulation train (200 Hz, 0.5 s, 2.5xT) that led to a very long lasting depolarization of proprioceptive axons (Tonic PAD, example in Fig. 5g) caused a facilitation of the monosynaptic EPSP that lasted for minutes (average shown, tonic cutaneous condition), and this was blocked by L655708 (in rats, n = 5 each). * P < 0.05: significant change with conditioning. + P < 0.05: significant change with antagonist. Raw data points shown to indicate occasional inhibition of the MSR by conditioning, but overall facilitation. ChR2 activation of GABAaxo neurons lacked these long tonic PAD-mediated after effects on the EPSP facilitation (Fig. 5e-f, Post), suggesting an additional source of GABA mediating after effects. f, Summary box plots of change in EPSP induced by cutaneous DR (DR1) conditioning (and associated phasic PAD) 60 ms prior to evoking the EPSP, with varying EPSP stimulation intensity. When the DR that evoked the test EPSP (DR2) was stimulated at an intensity that produced less than half the maximal EPSP height (1.1xT, \sim 30% max EPSP, n =42, same data as in e) the facilitation of EPSP by conditioning was larger than when this DR2 stimulation was increased to produce a test EPSP near maximal (1.5xT, prior to conditioning, n = 18). *P < 0.05: significant change with conditioning. This is likely because the stronger test stimulation reduced the headroom for increasing EPSPs by recruiting more proprioceptive axons, and increased self-facilitation prior to conditioning, the latter during repeated testing used to obtain EPSP averages. g, Summary of cutaneous facilitation of EPSPs from (f) (evoked by DR2 at 1.1xT), but separated into trials without (as in (b), n = 31) and with (as in (d), n = 11) large background postsynaptic changes induced by conditioning that lasted up to and during the EPSP testing (at 60 ms post conditioning, Bkg). *P < 0.05: significant change with conditioning evoked PAD. +P < 0.05: significant reduction facilitation with increased background activity (Bkg). h, Remote postsynaptic inhibition from conditioning. Long lasting changes in intrinsic proprieties of motoneurons (S4 and S3) following a mixed proprioceptive and cutaneous conditioning DR stimulation (on S3 or contralateral S4 DR, 2.5xT, DR1) that only produced a transient postsynaptic depolarization that ended prior to EPSP testing (as in B), including a reduction in time constant (τ) and slight hyperpolarization of potential (V_m), both measured at the time of EPSP testing (measured at 60 ms post conditioning, but in trials without EPSP testing; rats, n = 15). At this time, there was little change in somatic membrane resistance (Rm) with conditioning, suggesting that conditioning induced postsynaptic activity at a remote location in distal dendrites of the motoneuron. Indeed, when we voltage clamped the membrane potential during monosynaptic testing (DR2 at 1.1-1.5xT) to directly measure the synaptic current (EPSC) and minimize that inhibitory action of postsynaptic conductance increases, we found that the conditioning stimulation (DR1) produced a larger facilitation of the monosynaptic EPSC than the EPSP measured in current clamp in the same motoneurons. These results are consistent with the facilitation of the EPSP being masked by postsynaptic inhibition from increases in remote dendritic postsynaptic conductances triggered by the conditioning stimulation. *P < 0.05: significant change with conditioning.

Supplementary Table 1. Chronological list of evidence contradicting the classical concept of presynaptic inhibition of transmitter release from proprioceptive sensory axon terminals on motoneurons.

Date	Contradictions in classic view of terminal presynaptic inhibition mediated by terminal GABA _A receptors and PAD	Resolution of contradictions
1938	Primary afferent depolarization (PAD) directly evokes spikes in sensory axons, producing excitation rather than presynaptic inhibition. Barron and Matthews (1938) ²³ discovered that sensory nerve conditioning evokes a long depolarization in many other sensory afferents (primary afferent depolarization, PAD), which we now know is mostly GABA _A mediated ¹¹³ . They and others noted that sometimes this PAD was large enough to directly induce axon spiking, even in vivo ⁴⁶ , including spikes in the sensory axons mediating the MSR itself, raising a contradiction with the notion of GABA mediated presynaptic inhibition ¹⁶ . While these PAD-triggered spikes do not fully propagate antidromically out the DRs in many axons (they fail en route), they are actually initiated in most axons and more likely to conduct orthodromically ¹⁶ , making most axons and their motoneuron synapse refractory to subsequent testing (post activation depression). Indeed, numerous groups have shown that these spikes directly activate the MSR pathway ^{7,9,47,117} . Thus, these PAD-evoked spikes must inhibit afferent transmission in the MSR pathway by making axons refractory and producing post activation depression of their terminal synapse, even in humans where PAD evoked spikes occur ³⁶ . This indirect inhibition is GABA _A mediated and thus readily mistaken for presynaptic inhibition (sensitive to GABA _A antagonists) ^{118,119} . Even Eccles noted this issue, and showed that just the refractory period alone in the sensory axon inhibits the MSR ⁹ .	Post activation depression from PAD evoked spikes inhibits the MSR and masks facilitation of the MSR by nodal facilitation. We find that facilitation of the MSR by conditioning evoked PAD is always reduced when it is associated with a large enough conditioning stimulation to evoke spikes in sensory afferents, which likely results from post activation depression of axon transmission. This likely explains why Fink et al. 7,17 recently saw inhibition of the MSR with optogenetic or sensory activation of GABA _{axo} neurons. Crucially, when looking for MSR facilitation, avoiding these spikes and post activation depression requires using weak conditioning stimuli, unlike previous studies 19,18
1949	Post-tetanic potentiation (PTP) of the MSR increases sensory nerve conduction, but its mechanisms have remained elusive. Lloyd (1949) ¹²⁰ concluded that increasing conduction along sensory axons (not just terminals) contributed to the minutes of facilitation of the MSR seen after a high frequency nerve stimulation train (PTP). However, he supposed this might be due to hyperpolarization of the sensory axons, even though we now know that such trains depolarize axons via tonic PAD ¹⁶ . The tonic PAD from these bursts must overwhelm the hyperpolarization driven by Na-K pump activity ¹²¹ . Lloyd also concluded that PTP only occurred when the same nerve is used for the train (tetanus) as for testing the MSR. PAD is associated with a lowering of the threshold for activating spikes. Early on Wall (1958) ³¹ noted that a conditioning nerve stimulation that depolarized sensory axons (PAD) was associated with a lower threshold to extracellularly activate these axons. Subsequently this was assumed to be due to the action of terminal GABA _A receptors and presynaptic inhibition, and spike threshold changes were used to estimate the size of PAD ^{113,122} .	Repetitive nerve stimulation produces a tonic GABA _A mediated depolarization (PAD) of axons that facilitates nodal conduction, and increases the MSR. This PAD likely contributes to PTP, and is largest when the same nerve is tetanized, compared to other nerves, explaining why Lloyd missed the subtler facilitation from other nerves. PAD lowers the spike threshold via nodal GABA _A receptors assisting the sodium spike. This is not related to presynaptic inhibition, but can still be used to estimate PAD as Rudomin and others have done.
1957 - 1994	PAD is not correlated with inhibition of the monosynaptic reflex (MSR). Shortly after Frank and Fortes discovered that the leg extensor muscle MSR is inhibited by a conditioning of a flexor nerve in cats (PBST; like Fig. 6) ^{123,124} , Eccles proposed the concept of presynaptic inhibition mediated by this conditioning depolarizing of the proprioceptive sensory axon terminals in the MSR pathway (PAD), simply because the MSR and PAD are somewhat correlated in time ⁹ . However, in retrospect PAD is far too brief to account for the much longer inhibition caused by this conditioning ^{43,115} , and some flexor nerve conditioning (a single PBST pulse) inhibits the MSR (Fig. 1 of Eccles, 1961 ⁹), even though it does not cause PAD in the extensor proprioceptors of the MSR at all ¹²⁵ .	PAD is correlated with nodal spike facilitation and facilitation of the MSR. PAD causes facilitation of the MSR, explaining this correlation. When PAD is large and evokes axonal spikes, these cause post activation depression (detailed above) that should also be correlated with PAD, but is not due to presynaptic inhibition. Also, barbiturates used by Eccles and others potentiated GABAA receptor currents.
1959 - 1993	Postsynaptic inhibition inevitably accounts for part of the inhibition of the MSR by flexor nerve conditioning. In his initial short report Frank (1959) ¹²³ correctly suggested that the early inhibition of the MSR by flexor nerve conditioning might be partly postsynaptic (rather than presynaptic), on motoneuron distal dendrites. Others dismissed postsynaptic inhibition because the decay times of the EPSP does not always change when the EPSP is reduced by conditioning, which they proposed indicated that there was no postsynaptic change in conductance in distal dendrites ^{45,113} . However, this method is likely not very sensitive (due to variability in unitary EPSP time course). Also, anatomically ~70% of GABA _{axo}	Postsynaptic inhibition masks facilitation of the MSR by nodal facilitation. We find evidence for long lasting postsynaptic inhibition on distal motoneuron dendrites during nerve conditioning stimulation (including postsynaptic reductions in Tau, Vm and unitary EPSP heights and single MU firing). Crucially, minimizing

contacts on afferent terminals also contact motoneurons (in a triad), so postsynaptic inhibition requires using postsynaptic inhibition is inevitable 8,44. a small conditioning stimulation when looking for MSR facilitation, unlike previous studies 18. 1961 Self-facilitation during repeated MSR testing reduces the possibility of observing Self-facilitation masks facilitation of facilitation with subsequent conditioning stimuli, leaving only inhibitory actions of the MSR by a conditioning stimulation. 2014 conditioning. Eccles and others knew that the same proprioceptive nerve To observe facilitation of the MSR by a stimulation that activates the MSR also depolarizes these proprioceptive afferents conditioning stimuli that produces a (PAD self-activation)⁹. Thus, just the act of repeatedly testing the MSR to find the PAD it is important to use long test average MSR prior to conditioning pre-activates PAD and produces self-facilitation intervals (5 - 10 s) and small MSR test of the MSR, reducing the headroom to observe changes in the MSR following a intensities (1.1xT) to minimize self separate nerve conditioning stimulation that produces PAD. However, at the time activation of a tonic PAD prior to it was not known that repeated nerve stimulation causes a tonic buildup of GABA conditioning. While experimentally and a tonic PAD that alters sensory transmission and MSR even at long repetition troublesome, self facilitation during intervals of many seconds. Thus, Eccles and others used short test intervals (1 s) repetitive activation is actually one of and strong maximal MSR test stimuli 7,9,18, presumably assuming that there would the main functions of PAD, allowing be no interaction between test stimuli, which is not the case. In retrospect, these sensory axons to faithfully transmit short test intervals and strong test stimuli must have preactivated tonic GABA, spikes to motoneurons at high leaving little headroom to observe facilitation of the MSR (facilitation), and leaving frequencies that would otherwise mainly only inhibitory action possible. produce sodium spike inactivation. 1980 Sensory axon terminal potentials at motoneurons are consistent with spike failure. Positive terminal potential fields are Early efforts to examine how spikes propagated to sensory axon terminals decreased with PAD, indicative of employed extracellular recordings (EC) near the motoneurons, called terminal decreased conduction failure, and in potentials (Sypert et al. 1980)¹¹⁶. However, unlike EC recordings from near retrospect consistent with Sypert¹¹⁶. conducting axons (Fig. 2b), these terminal potentials lacked much of the obvious There is a small negative field that negative field associated with the action potential, and instead had a prominent follows the positive field in terminal positive field, followed by a smaller negative field (Extended Data Fig. 9 and potential recordings, representing Sypert¹¹⁶). This positive field has been shown in other axons to be indicative of spikes that actually reach the spike propagation failure and result from the passive axonal current caused by the terminals. We quantified negative field last non-failing node, similar to a FP, as demonstrated in motor axon recordings and found it to increase with PAD, ^{71,73}. Indeed, we found that even dorsal horn recordings could exhibit this positive consistent again with increased spikes field if the nearby dorsal root conduction is blocked with a microinjection of TTX conducting to motoneurons (Extended (Extended Data Fig. 9d). Sypert¹¹⁶ went on to show that with PAD evoked by nerve Data Fig. 9). Blocking activity in the conditioning this positive terminal potential field was decreased, and incorrectly spinal cord with glutamate interpreted this as evidence for decreased spike conduction and thus supposed it antagonists, which would include was due to presynaptic inhibition. blocking GABA_{axo} circuit activity, decreased this negative field. 1988 GABA_B receptors cause presynaptic inhibition and related RDD. Decades, after GABA_B mediated presynaptic inhibition Eccles popularized the notion of GABA_A mediated presynaptic inhibition, Curtis masks facilitation of the MSR by nodal (1998)^{43,118} concluded that the late part of the inhibition of the MSR by flexor 1998 GABA_A receptors. GABA_B receptors are nerve conditioning is instead GABA_B receptor mediated, since it is reduced by the located on the terminals, and produce GABA_B antagonist CGP55845 (as Fink also showed¹⁷), and as is RDD¹²⁶. RDD is a presynaptic inhibition (Fig. 5) and RDD rate dependent depression in the MSR during repeated testing. We suggest RDD is (Bennett and Hari, unpublished partly mediated by a build up of GABA released by GABAergic neurons onto the results), which are reduced by GABAB terminals during this repeated MSR testing, though activity dependent antagonists (CGP55845) or silencing homosynaptic depression likely also contributes¹²⁷. GABAaxo neurons. 1990 GABA_A receptors have direct postsynaptic inhibitory effects on many spinal GABA_A receptor antagonists reduce the neurons, making the actions of GABA_A antagonists difficult to attribute to MSR, by reducing nodal facilitation. 1998 presynaptic inhibition. By the 1990s Redman and others tried to confirm the role Postsynaptic GABA_A receptors have of GABA_A receptors in presynaptic inhibition by locally applying the GABA_A potent inhibitory actions on many antagonists bicuculline to the spinal cord, and indeed found this drug or other spinal neurons involved in antagonists reduced the inhibition of the MSR by flexor nerve conditioning polysynaptic reflexes. However, ^{18,43,118,128}. However, we now know that this is indirectly due to bicuculline causing minimizing these polysynaptic reflexes a widespread disinhibition of the spinal cord (including loss of post activation (by using weak test stimuli and depression, detailed above) that leads to a convulsive spinal cord with very long blocking NMDA receptors) reveals a lasting polysynaptic reflexes evoked by the nerve conditioning or the MSR testing direct inhibition of the MSR by GABAA itself, making pre and postsynaptic actions hard to distinguish. Further, we know antagonists, as does optogenetically

silencing GABA_{axo} neuron, consistent

with GABA_A receptors facilitating

rather than inhibiting sensory

transmission.

that GABA_A receptors mediate dorsal root reflexes and associated post activation

depression of the MSR (see above point), and thus bicuculline and picrotoxin likely

reduce the inhibition of the MSR via reducing post activation depression (see

above), rather than changing presynaptic inhibition.

1990	PAD recorded in the dorsal roots cannot arise from terminal GABA receptors, due	Space constant λ_s of sensory axons is
- 1995	to spatial attenuation on the axon. With advent of detailed anatomical and computer models of sensory axons ^{34,129,130} it became clear that signals like spikes	about 90 μm. Thus, the PAD recorded in the dorsal root must arise from
1995	or PAD are attenuated over short distances in axons, < 200 μm. This implies that	nearby nodal GABA receptors, and not
	PAD recorded on or near the DR is unlikely to bare any relation to terminal	bear any relation to GABA action at
	presynaptic inhibition, despite claims to the contrary ^{7,37,113,128} .	the terminals >1000 μm away.
1994	Shunting inhibition produced by axon terminal GABA _A receptors is not adequate to	GABA _A receptors only slightly decrease
-	produce presynaptic inhibition of the MSR. Numerous invertebrate studies	spikes by shunting conductances, and
1999	proposed that the conductance from GABA _A receptors in terminals caused a	otherwise assist nodal spike
	reduction in spike height via its shunting action that contributed to presynaptic	conduction in proprioceptive axons. In
	inhibition with nerve conditioning ²⁴ . However, the effects of conditioning on	non-failing secure spikes in sensory
	spikes was small and terminals were not actually recorded from. Subsequently	axons GABA _A receptors lower the
	modelling considerations led to the conclusion that shunting inhibition is not	threshold for spike activation
	adequate to produce presynaptic inhibition and calcium was somehow involved ¹²⁹ , possibly further implicating GABA _B receptors, as we see. Considering	(rheobase) and speed the spikes, the latter by decreasing the time constant
	our estimated space constant λ_S of ~90 µm, the small shunting inhibition of the	of the axon (RC). They do decrease the
	spike height (1 mV) we observe is very unlikely to prevent the spike produced at a	spike, but only by about 1%, consistent
	given node from activating a downstream neighbouring node, since nodes are ~50	with shunting being unlikely to inhibit
	μm apart, leading to only about a 50% reduction in spike height at the	spike transmission to motoneurons.
	downstream node (to ~40 mV, unless of course the node is failing), which is well	However, this does not rule out
	above that needed to initiate a full nodal spike. Thus, spike propagation is very	densely expressed GABA _A receptors
	unlikely to be blocked by shunting inhibition. Also, terminal boutons are mostly on	causing shunting and presynaptic
	unmyelinated axons without sodium channels (passive, 3rd order), and so a 1%	inhibition in cutaneous afferents, as
	reduction in the spike arising from the last/closest node on the 2nd order branch	previously suggested ^{16,70,131} .
	will have little effect on the terminal depolarization (1%), ruling out substantial shunting inhibition of transmitter release from the terminal.	
1994	Sodium spike inactivation from axon terminal GABA $_A$ receptor depolarization is not	Physiological PAD depolarizations do
-	adequate to produce presynaptic inhibition. Early poor quality recordings from	not block proprioceptive sensory axons
1998	sensory axons (resting near -50 mV from penetration injury) ¹³² led to the	spikes, and instead prevent them from
	prevailing view that spike failure with depolarization (PAD) was much more	failing in the MSR pathway. However,
	common than we now find with better recordings (resting near -70 mV, Extended	this does not rule out densely
	Data Fig. 3b). Further, Redman later questioned this view, and it seems unlikely	expressed GABA _A receptors causing
4005	for the MSR pathway ^{18,119} .	spike inactivation in other axons.
1995	Physiological GABA _A receptor activation is unlikely to produce branch point failure in the consequence of the MCP pathway. Over the years consequence are a superior of the MCP pathway.	GABA _A receptors help prevent branch
1998	in the sensory axons of the MSR pathway. Over the years sensory axon conduction failure has been occasionally noted from indirect observations ^{28,29,70,133-136} , and	point failure and thus facilitate sensory transmission in the MSR. Computer
1556	Wall and others ^{34,70,131} questioned whether this failure could be increased by	simulations by Walmsley and others
	GABA. However, Wall thought GABA should inhibit rather than assist spikes by	^{34,129} have led to the conclusion that
	inactivating sodium channels. However, Wall was misled by two issues. First, at	physiological GABA _A receptor
	the time poor quality recordings from sensory axons led to the misconception that	conductances cannot stop spikes from
	spike failure with physiological depolarizations (like PAD) was common ¹³² . To be	propagating past a node. Instead we
	fair, Wall was studying cutaneous, as well as proprioceptive, afferents, which are	report here that they help prevent
	more densely innervated by GABA receptors ¹⁶ , making spike inactivation by PAD	spike failure at branch points,
	more likely ⁷⁰ . Second, by this time GABA _A and associated PAD had been firmly entrenched as synonymous with presynaptic inhibition.	including in our computer simulations.
1996	Little or no terminal GABA _A receptors on proprioceptive axons. Extrasynaptic α 5	GABA _A receptors are mostly at nodes,
-	GABA _A receptors were found to be lacking at most proprioceptive axon terminals	whereas $GABA_B$ receptors are at
2018	in the ventral horn ¹⁶ . Synaptic GABA _A receptor labelling on these terminals had	terminals in large proprioceptive
	not been directly addressed, though see ^{6,7,137} . GABA _B receptor immunolabelling	afferents.
	had also not been investigated in these afferents, though is has in others $(A\beta)^{138}$.	
2005	When GAD2 expressing GABAergic neurons were identified that make axoaxonic	A key role of GABA _{axo} neurons is to
-	connections onto terminals of proprioceptive sensory axons ⁶⁻⁸ they were presumed	innervate proprioceptive afferent
2014	to mainly produce presynaptic inhibition. Thus, they were called GABA _{pre} neurons	nodes via GABA, receptors and ventral
	(also called P-bouton neurons), unfortunately implying that their only role is only presynaptic inhibition. We have instead referred to these neurons by the more	terminals via GABA _B receptors, producing nodal facilitation and
	generic termed GABA _{axo} .	producing nodal facilitation and presynaptic inhibition, respectively.
2018	GABA $_{0XO}$ neuron activation by sensory conditioning does not depolarize	$GABA_{axo}$ neuron activation depolarizes
	proprioceptive axon terminals. Direct recordings from the fine terminals of	nodes. Dorsally located nodes produce
	proprioceptive afferents reveal that during sensory conditioning the terminal is	the PAD recorded dorsal roots.
	not depolarized during the long PAD recorded on dorsal roots ¹⁶ .	

Supplementary Table 2. Resources used in Methods.

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Rabbit anti-α₅ GABA _A receptor subunit	OriGene Tech.	TA338505
Rabbit anti- α ₁ GABA _A receptor subunit	Sigma-Aldrich	06-868
Guinea pig anti-α₂ GABAA receptor subunit	Synaptic Systems	224 104
Chicken anti-γ ₂ GABA _A receptor subunit	Synaptic Systems	224 006
Rabbit anti-GABA _{B1} receptor subunit	Synaptic Systems	322 102
Mouse anti-NF200 (Neurofilament 200)	Sigma-Aldrich	N0142
Guinea pig anti-VGLUT1 (Vesicular glutamate transporter 1)	Sigma-Aldrich	AB5905
Rabbit anti-Caspr (Contactin associated protein)	Abcam	ab34151
Mouse anti-Caspr (Contactin associated protein)	NeuroMab	K65/35
Chicken anti-MBP (Myelin basic protein)	Abcam	ab106583
Chicken anti-VGAT (Vesicular inhibitory amino acid	Synaptic Systems	131 006
transporter)		
Rabbit anti-VGAT	Sigma-Aldrich	AB5062P
Rabbit anti-EYFP (Enhanced yellow fluorescent protein)	Biorbyt	orb256069
Goat anti-RFP (Red fluorescent protein; binds tdTom)	Biorbyt	orb334992
Rabbit anti-RFP (Red fluorescent protein; binds tdTom)	MBL Int.	PM005
Rabbit anti-GFP (Green fluorescent protein)	ThermoFisher Sc.	A11122
Mouse anti-Pan Sodium Channel (binds all Na _V types)	Sigma-Aldrich	S8809
Goat anti-rabbit Alexa Fluor 555	ThermoFisher Sc.	A32732
Goat anti-rabbit Alexa Fluor 647	Abcam	ab150079
Goat anti-rabbit Pacific orange	ThermoFisher Sc.	P31584
Goat anti-mouse Alexa Fluor 647	ThermoFisher Sc.	A21235
Goat anti-mouse Alexa Fluor 488	ThermoFisher Sc.	A11001
Goat anti-mouse Alexa Fluor 555	ThermoFisher Sc.	A28180
Goat anti-guinea pig Alexa Fluor 647	ThermoFisher Sc.	A21450
Goat anti-chicken Alexa Fluor 405	Abcam	ab175674
Donkey anti-goat Alexa Fluor 555	Abcam	ab150130
Donkey anti-rabbit Alexa Fluor 488	ThermoFisher Sc.	A21206
Streptavidin-conjugated Alexa Fluor 488	Jackson immunoR.	016-540-084
Streptavidin-conjugated Cyanine Cy5	Jackson immunoR.	016-170-084
Chemicals, Peptides, and Recombinant Proteins		
M.O.M (Mouse on Mouse Immunodetection Kit)	Vector	BMK-2202
Experimental Models: Organisms/Strains		
Gad2 ^{CreER} mouse:	The Jackson Laboratory	Stock# 010702
Gad2 ^{tm1(cre/ERT2)Zjh} /J		
Vglut1 ^{cre} mouse:	The Jackson Laboratory	Stock# 023527
B6;129S-S/c17a7 ^{tm1.1(cre)Hze} /J		
R26 ^{LSL-ChR2-EYFP} mouse:	The Jackson Laboratory	Stock# 012569
B6;129S-Gt(ROSA)26Sor ^{tm32(CAG-COP4*H134R/EYFP)Hze} /J	T	Ct. 1 !! 042725
R26 ^{LSL-Arch3-GFP} mouse:	The Jackson Laboratory	Stock# 012735
B6;129S-Gt(ROSA)26Sor ^{tm35.1} (CAG-aop3/GFP)Hze/J	The legisles is the section	C+0.0k# 00701 #
R26 ^{LSL-tdTom} mouse crossed with Gad2 ^{CreER} mice: B6.Cg- <i>Gt(ROSA)26Sor</i> ^{tm14(CAG-tdTomato)Hze} /J	The Jackson Laboratory	Stock# 007914
R26 ^{LSL-tdTom} mouse crossed with Vglut1 ^{Cre} mice:	The lackson Laboratory	Stock# 007909
B6.Cg-Gt(ROSA)26Sor ^{tm9(CAG-tdTomato)} Hze/J	The Jackson Laboratory	310CK# 00/303
Bolog Ottinophizoon		

Oligonucleotides		
5' -> ACG TTT CCT GTC CCT GTG TG -> 3'	Integrated DNA	11400
Common for Gad2 ^{CreER} mice	technologies	
5' -> AGG CAA ATT TTG GTG TAC GG -> 3'	Integrated DNA	oIMR9074
Mutant for Gad2 ^{CreER} mice	technologies	
5' -> CAG ACG CTG CAG TCT TTC AG -> 3'	Integrated DNA	oIMR3346
Wild type for Gad2 ^{CreER} mice	technologies	
5' -> ACA TGG TCC TGC TGG AGT TC -> 3'	Integrated DNA	oIMR9102
Mutant Forward for ChR2 mice	technologies	
5' -> GGC ATT AAA GCA GCG TAT CC -> 3'	Integrated DNA	oIMR9103
Mutant Reverse for ChR2 mice	technologies	
5' -> AAG GGA GCT GCA GTG GAG TA -> 3'	Integrated DNA	oIMR9020
Wild type Forward for ChR2 mice	technologies	
5' -> CCG AAA ATC TGT GGG AAG TC -> 3'	Integrated DNA	oIMR9021
Wild type Reverse for ChR2 mice	technologies	
5' -> CTG TTC CTG TAC GGC ATG G -> 3'	Integrated DNA	oIMR9105
Mutant Forward for both tdTom mouse strains	technologies	
5' -> GGC ATT AAA GCA GCG TAT CC -> 3'	Integrated DNA	oIMR9103
Mutant Reverse for both tdTom mouse strains	technologies	
5' -> AAG GGA GCT GCA GTG GAG TA -> 3'	Integrated DNA	oIMR9020
Wild type Forward for both tdTom mouse strains	technologies	
5' -> CCG AAA ATC TGT GGG AAG TC -> 3'	Integrated DNA	oIMR9021
Wild type Reverse for both tdTom mouse strains	technologies	
5' -> CTT CTC GCT AAG GTG GAT CG -> 3'	Integrated DNA	12178
Mutant Forward for Arch3 mice	technologies	
5' -> CAC CAA GAC CAG AGC TGT CA -> 3'	Integrated DNA	12179
Mutant Reverse for Arch3 mice	technologies	
5' -> TCC CAA AGT CGC TCT GAG TT -> 3'	Integrated DNA	oIMR8713
Wild type Forward for Arch3 mice	technologies	
5' -> CTT TAA GCC TGC CCA GAA GA -> 3'	Integrated DNA	12177
Wild type Reverse for Arch3 mice	technologies	
5' -> ATG AGC GAG GAG AAG TGT GG -> 3'	Integrated DNA	17904
Common for VGLUT1 ^{cre} mice	technologies	
5' -> CCC TAG GAA TGC TCG TCA AG -> 3'	Integrated DNA	12231
Mutant reverse for VGLUT1 ^{cre} mice	technologies	
5' -> GTG GAA GTC CTG GAA ACT GC -> 3'	Integrated DNA	17905
Wild type reverse for VGLUT1 ^{cre} mice	technologies	

Software and Algorithms		
Leica Application Suite X software	Leica Microsystems	
Clampfit 8.0	Axon Instruments	