Activation of cortical somatostatin interneurons prevents the development of neuropathic pain

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Neuropathic pain involves long-lasting modifications of pain pathways that result in abnormal cortical activity. How cortical circuits are altered and contribute to the intense sensation associated with allodynia is unclear. Here we report a persistent elevation of layer V pyramidal neuron activity in the somatosensory cortex of a mouse model of neuropathic pain. This enhanced pyramidal neuron activity was caused in part by increases of synaptic activity and NMDA-receptor-dependent calcium spikes in apical tuft dendrites. Furthermore, local inhibitory interneuron networks shifted their activity in favor of pyramidal neuron hyperactivity: somatostatin-expressing and parvalbumin-expressing inhibitory neurons reduced their activity, whereas vasoactive intestinal polypeptide–expressing interneurons increased their activity. Pharmacogenetic activation of somatostatin-expressing cells reduced pyramidal neuron hyperactivity and reversed mechanical allodynia. These findings reveal cortical circuit changes that arise during the development of neuropathic pain and identify the activation of specific cortical interneurons as therapeutic targets for chronic pain treatment.

Neuropathic pain is a chronic pain state resulting from peripheral and/or central nerve injury1. The development of persistent pain is believed to be due to the long-lasting changes of neuronal functions in the pain transmission pathway, from peripheral nociceptors and the spinal cord to supraspinal and cortical areas including the primary somatosensory cortex (S1) and the anterior cingulate cortex (ACC)2–6. S1 is an important cortical area responsible for the sensory aspects of the pain, such as the intensity and location of the pain7,8. This cortical region is consistently activated during nociception and becomes hyperactive under chronic pain states. In both human and animal studies, S1 of nerve-injured subjects shows increased activation, somatotopic reorganization and changes in cortical thickness9–11. Furthermore, strategies to reduce S1 hyperexcitability and reorganization demonstrate benefits against chronic pain development12–14.

How neuronal circuitry in S1 is modified and becomes hyperactive during the development of neuropathic pain remains unclear. At the cellular level, chronic pain states have been associated with structural and functional alterations in cortical pyramidal neurons, including long-term potentiation of synaptic transmission, new dendritic spine formation, increases in intrinsic cellular excitability and rearrangements in ion channels15–20. In addition to these changes in glutamatergic pyramidal neurons, alterations of a highly interconnected network of GABAergic interneurons may also contribute to the persistent pain state. For example, increased activity of both excitatory neurons and local interneurons in layer II/III (L2/3) of S1 was observed in a model of inflammatory pain21. Within S1, there is a large diversity of cortical interneurons, including parvalbumin (PV), somatostatin (SOM) and vasoactive intestinal polypeptide (VIP)–expressing neurons22. These different types of interneurons target specific domains of principal neurons or other interneurons, providing precise spatiotemporal control of excitatory and inhibitory outputs and cortical dynamics. How changes in glutamatergic pyramidal neurons and interneuron populations contribute to the hyperactivity in S1 and the intense sensation in neuropathic pain are unknown.

In this study, we investigated changes of layer V (L5) pyramidal neurons and three interneuron types in S1 after spared nerve injury (SNI), a model of neuropathic pain in mice23,24. By imaging and manipulating neuronal activity in living mice, we identified cell-type-specific changes in S1 that were important for the development of neuropathic pain. Our results also suggest modulating interneuron activity as an effective strategy for correcting pyramidal neuron hyperactivity and mitigating mechanical allodynia.

RESULTS

Peripheral nerve injury causes pyramidal neuron hyperactivity in S1

To identify cortical changes related to neuropathic pain, we used in vivo two-photon calcium imaging to examine somatic activity of pyramidal neurons expressing the genetically encoded Ca2+ indicator GCaMP6s in S1 of awake, head-restrained mice. To induce persistent neuropathic pain, mice were subjected to SNI (see Online Methods)23,24 (Fig. 1a). Two days after SNI, mice experienced mechanical allodynia in the injured hind paw, as revealed by a marked reduction in the paw withdrawal threshold upon pressure application to the lateral aspect of the plantar paw surface by a von Frey filament (Fig. 1b). This mechanical hypersensitivity persisted for more than 1 month. In contrast, sham-operated mice did not exhibit pain behavior when assessed after 1 week. Both SNI and sham mice showed no changes in paw withdrawal threshold in intact limbs.

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We first examined the activity of L5 pyramidal neurons, the major output cells in S1 that integrate sensory inputs from thalamic and intracortical origin. In the region of S1 corresponding to the injured hind limbs of the animals, we found that Ca2+ activity in the somata of L5 pyramidal neurons (located >600 µm below the pial surface) but not ipsilateral, was apparent, and lasted for at least 2 months. Moreover, intraperitoneal injection of MK801, an antagonist of NMDA receptors, reduced the level of L5 Ca2+ activity in SNI mice to that in sham mice 1 month after surgery.

Fig. 1a). We performed an increase of Ca2+ activity in S1 after peripheral nerve injury. (a) Schematic of SNI and two-photon Ca2+ imaging timeline in S1. Imaging planes were different cortical depths (dashed boxes) from the pial surface in awake, head-restrained mice expressing GCaMP6s in L5 pyramidal (PYR) neurons. (b) Measures of hind limb paw withdrawal threshold before and after SNI (n = 17 mice) and sham operations (n = 12 mice) (2 d: P = 0.025; 7 d: P < 0.001; 14 d: P < 0.001; 31 d: P < 0.001, two-way ANOVA followed by Tukey’s test). (c) Fluorescence traces from representative L5 PYR somata expressing GCaMP6s in SNI or sham mice 1 month after surgery. (d) Average total integrated Ca2+ activity over 2.5 min in L5 PYR 1 month after surgery (SNI: 116.2 ± 8.5 ΔF, n = 141 cells from 5 mice; sham: 41.6 ± 2.9 ΔF, n = 59 cells from 5 mice; t198 = 5.581, P < 0.001, unpaired t test; SNI ipsilateral: 20.2 ± 1.0 ΔF, n = 106 cells, t25 = 9.694, P < 0.001, unpaired t test). (e) L5 PYR soma Ca2+ activity at 1 month correlates with the hind limb paw withdrawal threshold in SNI mice (Pearson r = −0.33, P < 0.001), but not in sham mice (Pearson r = 0.046, P = 0.609). (f) L5 PYR soma total integrated Ca2+ activity before (SNI: 40.3 ± 3.6 ΔF, n = 50 cells from 2 mice; sham: 17.1 ± 1.2 ΔF, n = 45 cells) and after local application of TTX to sciatic nerve (SNI: 41.9 ± 3.7 ΔF, t49 = 1.928, P = 0.06, paired t test; sham: 11.6 ± 0.9 ΔF, t49 = 12.08, P < 0.001, paired t test from 2 mice) 2 weeks after sham surgery or SNI. Data are presented as means ± s.e.m. *P < 0.05, ***P < 0.001. (c) Representative traces from experiments carried out on at least 5 animals per group.

In addition to hyperactivity of L5 pyramidal neurons, we found that the somata of L2/3 pyramidal neurons (located 200–300 µm below the pial surface) also exhibited higher levels of Ca2+ activity in SNI mice as compared to sham mice (Supplementary Fig. 2). As in L5 neurons, this elevated neuronal activity could be observed shortly after SNI, at 1 week, and was sustained over 2 months (Supplementary Fig. 2). Together these findings indicate that peripheral nerve injury causes a robust and persistent hyperactivity of pyramidal neurons in S1.

**Increased dendritic spine and branch activity in mice with neuropathic pain**

The increased activity of L5 pyramidal neurons associated with neuropathic pain could be due to either an increase of excitatory drive onto these neurons, a reduction of inhibition, enhanced intrinsic excitability,
Figure 2 Enhanced dendritic spine and branch activity in S1 of neuropathic pain mice. (a) Representative two-photon images of active, apical tuft dendrites of L5 PYR neurons expressing GCaMP6s in SNI or sham mice 1 month after surgery. Arrowheads point to spine heads measured. (b) Fluorescence traces of dendritic spines and adjacent shaft show increased Ca\(^{2+}\) transients in spines and shaft of SNI as compared to sham mice. Light magenta bars indicate Ca\(^{2+}\) elevations in both spine heads and shaft, indicating Ca\(^{2+}\) spike generation. (c) Distribution of peak Ca\(^{2+}\) amplitude of dendritic spines observed in both SNI and sham mice. Notably, we found that SNI mice showed a marked increase in dendritic spine Ca\(^{2+}\) activity as compared to sham mice (Fig. 2a–d). Furthermore, spine pairs randomly selected from individual dendritic branches displayed greater coactivity in SNI mice than in sham mice (Fig. 2e,f).
2+ activity in spines, we also observed 2+ spike generation in apical tuft branches and trunks of L5 pyramidal neurons in S1 of neuropathic pain mice. Fullerenes, we also observed 2+ spike generation in apical tuft branches and trunks of L5 pyramidal neurons in S1 of neuropathic pain mice.

In addition to localized Ca2+ activity in spines, we also observed Ca2+ transients generating across long segments (>30 µm) of L5 apical tuft dendrites. These dendritic Ca2+ transients occurred across long stretches of dendrites with comparable ΔF/ΔF0 and lasted several seconds (Fig. 2a,g). They were similar to the dendritic Ca2+ spikes observed in pyramidal neurons in other cortical regions of living animals26,27. When comparing SNI to sham mice, we found that dendritic Ca2+ spikes occurred more frequently in SNI mice (Fig. 2a,h). Furthermore, the average peak amplitude (Fig. 2i) and average total integrated activity (Fig. 2j) of these dendritic Ca2+ spikes were also higher in SNI mice than in sham mice.

Like the somatic activity, enhanced tuft Ca2+ activity of L5 pyramidal neurons persisted from 1 week to 2 months (Supplementary Fig. 3). When comparing SNI to sham mice, we found that dendritic Ca2+ activity on multiple sibling branches of L5 pyramidal neurons depends on synaptic activity originating in L1. Collectively, these results indicate that, following the induction of neuropathic pain, a substantial fraction of apical tuft dendrites of L5 pyramidal neurons in S1 receive synchronized synaptic inputs and generate persistent, NMDA-receptor-dependent Ca2+ spikes, which lead to the hyperactivity of L5 pyramidal neurons.

**Figure 3** Elevated dendritic Ca2+ spike generation in apical tuft branches and trunks of L5 pyramidal neurons in S1 of neuropathic pain mice. (a) Two-dimensional projection of sibling branches from an individual L5 PYR neuron in SNI mice. Five regions of interest (ROIs) (magenta boxes) corresponding to different dendritic branches were analyzed. All ROIs generated multiple Ca2+ transients during recording. (b) Percentages of local or global Ca2+ spikes observed on sibling branches from individual PYR neurons (SNI, 9.8 ± 1.2 transients/min, n = 24 cells from 3 mice; sham, 4.8 ± 1.0 transients/min, n = 16 cells from 3 mice). Ca2+ events were analyzed over 2.5 min. (c) Fluorescence traces of apical dendritic trunks imaged ~300 µm below the pia in SNI or sham mice. (d) Average total integrated Ca2+ activity over 2.5 min in apical trunk (SNI, 77.2 ± 4.8 ∆F, n = 102 trunks from 5 mice; sham, 32.9 ± 2.3 ∆F, n = 103 trunks from 5 mice; t255 = 9.34, P < 0.001). Local application of MK801 in SNI (16.7 ± 0.8 ∆F, n = 26 trunks from 2 mice, t255 = 8.111, P < 0.001, two-way ANOVA followed by Bonferroni’s test) and sham mice (8.3 ± 0.9 ∆F, n = 28 trunks from 2 mice, t255 = 3.4, P = 0.0016). (e) TTX locally applied to L1 (blue shaded circle) significantly reduced Ca2+ activity in apical trunk and L5 soma (noted by gray dashed lines) in SNI mice (before TTX: 1; trunk after TTX: 0.51 ± 0.02, n = 50 trunks, t49 = 23.77, P < 0.001, paired t-test; soma after TTX: 0.53 ± 0.03, n = 54 cells, t53 = 17.19, P < 0.001; trunks and somata chosen from different cells). Data are presented as means ± s.e.m. **P < 0.01, ***P < 0.001.

**Apical tuft Ca2+ transients contribute to L5 pyramidal cell hyperactivity**

Previous studies have shown that apical tuft dendrites exhibit Ca2+ spikes in either entire or a subset of branches in various cortical regions26,27. These dendritic Ca2+ spikes substantially boost the firing of L5 pyramidal neurons26,28. To better understand the contribution of apical tuft activity to the hyperactivity of L5 pyramidal neurons, we performed Ca2+ imaging at different cortical depths from the pial surface. In higher-order tuft branches converging toward the nexus (the primary branch point from the apical trunk, located 150–250 µm from the pial surface), we observed elevated dendritic Ca2+ activity on multiple sibling branches of L5 pyramidal neurons (Fig. 3a). At this cortical depth, the majority of Ca2+ transients (sham, 58 ± 4.1%; SNI, 67 ± 6.2%) occurred simultaneously in all branches (that is, were global) of the same L5 pyramidal neuron (Fig. 3b). At the apical trunk nexus of L5 pyramidal neurons, located 250–350 µm below the pial surface, repetitive Ca2+ activity occurred in both SNI and sham mice (Fig. 3c). The total integrated trunk Ca2+ activity was greater in SNI than in sham mice 1 month after surgery (Fig. 3d).

We found that local application of MK801 to L1 reduced the apical trunk Ca2+ activity in both SNI and sham mice (Fig. 3d). Furthermore, local application of TTX to L1 reduced Ca2+ activity in both trunks and somata of L5 pyramidal neurons in SNI mice (Fig. 3e), suggesting that the apical trunk and somatic hyperactivity in L5 pyramidal neurons depends on synaptic activity originating in L1. Collectively, these results indicate that, following the induction of neuropathic pain, a substantial fraction of apical tuft dendrites of L5 pyramidal neurons in S1 receive synchronized synaptic inputs and generate persistent, NMDA-receptor-dependent Ca2+ spikes, which lead to the hyperactivity of L5 pyramidal neurons.
SOM and PV interneurons reduce activity while VIP cells increase activity in neuropathic pain

To further understand mechanisms underlying the hyperactivity of L5 pyramidal neurons in S1, we examined the activity of local inhibitory cells during the development of neuropathic pain. Recent studies have shown that SOM-positive inhibitory interneurons are important in regulating dendritic and somatic activity of pyramidal neurons\(^{27,29-31}\). To measure the impact of neuropathic pain on SOM cell responses, we used \textit{in vivo} two-photon \(Ca^{2+}\) imaging to examine the activity of SOM cells expressing GCaMP6s following SNI or sham surgeries. In this experiment, SOM-IRES-Cre mice were injected with a Cre-dependent adeno-associated virus (AAV) to induce the expression of GCaMP6s specifically in SOM cells. In contrast to the increased activity of pyramidal neurons, we found that SOM cell activity was reduced by half (52%) 1 month after SNI (Fig. 4a, b and Supplementary Fig. 4). This reduction in SOM cell activity was also observed in their axon fibers projecting to L1 (Fig. 4c, d). Furthermore, SOM cells imaged 1 week after surgery showed a reduction in neuronal activity in SNI as compared to sham mice (Supplementary Fig. 4). These \textit{in vivo} \(Ca^{2+}\) recordings show that peripheral nerve injury induces a potent and persistent reduction in SOM cell firing rates in S1.
In addition to SOM neurons, we also investigated whether PV-positive interneurons were affected by neuropathic pain using PV-IRES-Cre mice expressing GCaMP6s. We found that PV cells also displayed a significant reduction in neuronal activity in SNI mice as compared to sham mice 1 week and 1 month after surgery (Fig. 4e,f and Supplementary Fig. 5). The reduction (~30%) in PV cell activity at 1 month was not as robust as that of SOM cells (~50%) when comparing SNI to sham mice. Because SOM and PV neurons target different regions of pyramidal cells (tuft dendrites versus the perisomatic region), reduced activity from these cells...

**Figure 5** Acute activation of SOM neurons in S1 reduces L5 pyramidal neuron activity in mice with neuropathic pain. (a) Representative fluorescence traces of SOM neurons expressing GCaMP6s and hM3Dq-cre before and after CNO injection in SNI mice. (b) Percentage change in average total integrated Ca²⁺ activity of SOM somata over 2.5 min following CNO injection in SNI (382 ± 78%, n = 93 cells from 4 mice) and sham mice (361 ± 114%, n = 61 cells from 5 mice). SOM Ca²⁺ activity significantly decreased from baseline activity upon CNO injection (SNI: t_{25} = 4.912, P < 0.001; sham: t_{20} = 3.168, P < 0.01, paired t test). No significant difference was found between SNI and sham mice after CNO injection (t_{25.2} = 0.163, P = 0.87, unpaired t test). (c) Representative fluorescence traces of apical dendrites of L5 PYR neurons expressing GCaMP6s before and after CNO injection to activate SOM hM3Dq-positive neurons in SNI mice. (d) Percentage change in dendritic Ca²⁺ activity of L5 PYR neurons following CNO injection in SNI (−14.5 ± 5.5%, n = 29 dendrites from 3 mice; t_{19} = 2.522, P = 0.0176, paired t test) and sham mice (−11.5 ± 5.3%, n = 43 dendrites from 4 mice; t_{42} = 2.149, P = 0.0374). (e) Representative fluorescence traces of L5 PYR soma expressing GCaMP6s before and after CNO injection to activate SOM hM3Dq-positive neurons in SNI mice. (f) Percentage change in L5 PYR somatic Ca²⁺ activity following CNO injection in SNI (−17.9 ± 4.0%, n = 94 cells; t_{63} = 4.449, P < 0.001, paired t test) and sham mice (−13.7 ± 5.9%, n = 87 cells; t_{86} = 2.310, P < 0.05). (g) Percentage change in average total integrated Ca²⁺ activity of SOM somata following CNO injection in naive mice infected with hM3Di (−25.4 ± 10.6%, n = 20 cells from 2 mice). SOM Ca²⁺ activity significantly decreased from baseline upon CNO injection (t_{18} = 2.391, P = 0.03, paired t test). (h) Inactivating SOM cells with hM3Di exacerbated Ca²⁺ activity of L5 PYR dendrites (b, percentage change in L5 dendritic Ca²⁺ activity after CNO: 32.9 ± 14.1%, n = 60 dendrites, t_{59} = 2.35, P = 0.02, paired t test) and somata (i, percentage change in L5 PYR somatic Ca²⁺ activity after CNO: 31.7 ± 5.0%, n = 60 somata, t_{59} = 6.289, P < 0.001, paired t test) in naive mice. (j) Design for determining mechanical allodynia before and after CNO-induced SOM cell activation or inactivation. Thresholds measured before and 20 min after CNO injection in SNI or sham mice. vF, von Frey. (k, l) Withdrawal threshold after SOM activation by CNO in SNI (before CNO: 17.1 ± 0.1 g, after 20 min CNO: 4.0 ± 0.2 g, n = 14 mice; t_{13} = 12.97, P < 0.001, paired t test) and sham mice (before CNO: 4.8 ± 0.4 g, after 20 min CNO: 5.2 ± 0.2 g, n = 14 mice; t_{13} = 1.685, P = 0.12, paired t test). (m, n) Withdrawal threshold before and after CNO injection in naive mice infected with hM3Di. Inactivating SOM cells had no significant effects on the animals’ withdrawal threshold (contralateral paw to hM3Di-infected S1 P = 0.12, paired t test; ipsilateral paw: P = 0.13, paired t test). (m) Percentage change in average total integrated Ca²⁺ activity of PV somata following CNO injection in SNI (185 ± 29.7%, n = 81 cells from 3 mice) and sham mice (131 ± 24.5%, n = 48 cells from 3 mice). PV Ca²⁺ activity significantly increased from baseline activity upon CNO injection (SNI: t_{30} = 6.243, P < 0.001; sham: t_{20} = 5.339, P < 0.001, paired t test). No significant difference was found between SNI and sham mice after CNO injection (t_{20} = 1.262, P = 0.21, unpaired t test). (n) Withdrawal threshold after PV activation by CNO in sham (before CNO: 5.4 ± 0.3 g, after 20 min CNO: 4.5 ± 0.4 g, n = 8 mice; t_{7} = 2.305, P = 0.06, paired t test) and SNI mice (before CNO: 1.9 ± 0.3 g, after 20 min CNO: 2.1 ± 0.1 g, n = 8 mice; t_{7} = 0.7693, P = 0.47, paired t test). Data are presented as means ± s.e.m. *P < 0.05, **P < 0.01, ***P < 0.001.

(a,c,e) Representative traces from experiments carried out on at least 3 animals per group.
suggests a lack of inhibitory tone across the somatodendritic axis of pyramidal neurons to promote pyramidal cell hyperactivity in neuropathic pain.

In addition to SOM and PV, VIP-expressing interneurons constitute another major subtype of neocortical interneurons and are known to directly inhibit the activity of both SOM- and PV-expressing

Figure 6 Daily activation of SOM neurons following peripheral nerve injury prevents the development of neuropathic pain. (a) Top panel, experimental design for daily activation of SOM neurons following SNI or sham surgeries (CNO, orange syringe; vF, von Frey). For control, SNI and sham mice received saline injections. Bottom panel, cartoon of two-photon imaging planes corresponding to SOM somata (b), pyramidal somata (c,d) and pyramidal tuft dendrites (e,f). (b) Percentage change in average total integrated Ca\(^2+\) activity of SOM somata over 2.5 min following single CNO injection (inj.) in SNI (n = 23 cells from 2 mice) and sham mice (n = 26 cells from 2 mice). CNO injection significantly increased from baseline the activity of SOM cells for at least 12 h (P < 0.001) but not 24 h (sham, P = 0.19; SNI, P = 0.29). (c) Fluorescence traces of L5 PYR neuron somata expressing GCaMP6s after 1 week of daily CNO activation of SOM cells in SNI or sham mice. (d) Average total integrated Ca\(^2+\) activity over 2.5 min recording of L5 PYR somata following 1 week of daily SOM activation in SNI (14.4 ± 0.7 ∆F, n = 170 somata from 5 mice) and sham mice (14.4 ± 1.2 ∆F, n = 54 somata from 5 mice). No significant difference between SNI and sham mice treated with CNO (t\(_{279} = 0.7357, P = 0.9251\)).

(e) Fluorescence traces of L5 apical tuft dendrites expressing GCaMP6s after 1 week of daily SOM activation in SNI and sham mice. (f) Average total integrated Ca\(^2+\) activity over 2.5 min recording of L5 dendrites following 1 week of daily SOM activation in SNI (14.1 ± 0.7 ∆F, n = 159 dendrites from 5 mice) and sham mice (12.9 ± 0.8 ∆F, n = 37 dendrites from 5 mice). No significant difference between SNI and sham mice treated with CNO (t\(_{54} = 0.93, P = 0.362\)).

(g) Withdrawal threshold under various conditions. Daily SOM activation significantly increased withdrawal threshold in SNI mice as compared to SNI mice injected with saline (P < 0.05 at day 5 and P < 0.001 at days 7, 14, 21 and 28). Data are presented as means ± s.e.m. n.s., not significant; *P < 0.05, **P < 0.01, ***P < 0.001, paired t test in b, two-way ANOVA followed by Bonferroni’s test in d, f and Tukey’s test in g. (c,e) Representative traces from experiments carried out on at least 3 animals per group.
interneurons in vitro and in vivo. This connectivity scheme of VIP → SOM/PV → pyramidal cells is thought to amplify and modulate local neuronal activity. To determine whether VIP neurons might be involved in the alterations of SOM and PV cell activity in neuropathic pain, we infected VIP-ires-Cre mice with AAV encoding GCaMP6s and imaged Ca²⁺ activity of VIP neurons in SNI and sham mice. Consistent with the disinhibition hypothesis, we found a ~90% increase in the activity of VIP neurons in SNI as compared to sham mice 1 month after surgery (Fig. 4g,h). The increased VIP neuron activity was also observed 1 week after SNI as compared to the sham control (Supplementary Fig. 5). Taken together, these experiments indicate that the development of neuropathic pain is accompanied by distinct and persistent changes of neuronal activity in different types of interneurons, including reduced neuronal activity of SOM and PV cells and an activation of VIP-expressing cells (see model in Supplementary Fig. 5).

Transient activation of SOM interneurons dampens L5 pyramidal cell hyperactivity in neuropathic pain

Previous studies have shown that SOM cells regulate branch-specific Ca²⁺ spike generation in vivo and control electrical and biochemical signaling in dendritic spines. To test whether the reduction of SOM cell activity in SNI mice contributes significantly to the hyperactivity of L5 pyramidal neurons, we activated SOM cells under the neuropathic pain condition. To increase SOM cell activity in vivo, we specifically infected SOM cells of S1 with an AAV encoding Cre-dependent hM3Dq DREADD receptors (designer receptors exclusively activated by designer drugs) in SOM-Cre mice (Supplementary Fig. 6). Binding of the ligand clozapine N-oxide (CNO) to hM3Dq receptors activates Gq-coupled signaling, leading to membrane depolarization through inhibition of KCNQ channels, which increases neuronal firing in target cells. Indeed, CNO delivered by intraperitoneal injection to mice expressing hM3Dq specifically in SOM cells induced a more than threefold increase in SOM somatic Ca²⁺ activity in vivo in both SNI and sham mice (Fig. 5a,b). To determine the effect of transient activation of SOM cells on pyramidal neuron activity in neuropathic pain, we imaged GCaMP6s-expressing L5 pyramidal neurons in mice expressing hM3Dq in SOM cells. Ca²⁺ activity in dendrites and soma of L5 pyramidal neurons was measured before and 20 min after CNO injection. In both SNI and sham, we found that acute activation of SOM cells induced a reduction in total integrated Ca²⁺ activity in both apical tuft dendrites and somata of L5 pyramidal neurons (Fig. 5c–f). Moreover, the somatic Ca²⁺ activity of L2/3 pyramidal neurons was also decreased after acute SOM activation with CNO (Supplementary Fig. 7). In contrast to the reduction in L5 neuron activity induced by SOM-hM3Dq, inactivating SOM cells with the inhibitory DREADD variant Gc-coupled hM4Di exacerbated Ca²⁺ activity of L5 dendrites and somata in SNI and sham mice (Fig. 5g–i).

As a control, we infected SOM-Cre mice with a control AAV vector (TurboRFP). As expected, administration of CNO to mice that did not express hM3Dq receptors had no effect on L5 pyramidal neuron activity (Supplementary Fig. 8).

When we tested mechanical hypersensitivity in SOM-Cre mice infected with AAV-hM3Dq (Fig. 5j), we found that acute activation of SOM cells with a single injection of CNO significantly increased the plantar pressure required to elicit paw withdrawal in SNI mice, whereas CNO injection in sham mice exerted no effects on the animals’ behavior (Fig. 5k). SNI and sham mice infected with AAV-CMV-TurboRFP showed no changes in pain threshold after receiving a CNO injection (Supplementary Fig. 8). Furthermore, unlike the substantial reduction (52%) in SOM activity and reduced paw withdrawal threshold in SNI mice, we found that a milder reduction (25%) of SOM cell activity with AAV-hM4Di and CNO in naive mice had no effect on the animals’ paw withdrawal threshold (Fig. 5i). Taken together, these findings indicate that acute activation of SOM cells in S1 is sufficient to reduce pyramidal cell activity and improve mechanical allodynia in neuropathic pain.

In contrast to its effect on the activity of SOM cells, CNO treatment had milder effect (less than a twofold increase) on the neuronal activity of PV cells expressing hM3Dq in both SNI and sham mice (Fig. 5m and Supplementary Fig. 9). When we tested withdrawal threshold in PV-Cre mice infected with AAV-hM3Dq, CNO injection had no effects on the animals’ pain sensation in either SNI or sham mice (Fig. 5n), suggesting that increasing PV cell activity with the DREADD-CNO approach is less effective at alleviating mechanical allodynia than increasing SOM cell activity.

Continuous SOM cell activation prevents the development of mechanical allodynia

The results above highlight the immediate beneficial effects of SOM interneuron activation in neuropathic pain at the cellular and behavioral level. However, it remains unknown whether continuous activation of SOM cells might preclude the establishment of mechanical allodynia. To test this possibility, following the SNI or sham surgery in mice expressing hM3Dq in SOM cells, we performed daily CNO injections for 1 week and imaged Ca²⁺ activity in L5 pyramidal neurons 1 d after the last CNO treatment (Fig. 6a). Consistent with previous reports, a single injection of CNO activated SOM cells over a period of ~12 h (Fig. 6b). We found that daily CNO treatment for 1 week was sufficient to prevent SNI-induced pyramidal neuron hyperactivity (Fig. 6c,d). At the level of apical tuft dendrites, we found no significant difference in the average Ca²⁺ activity between SNI and sham mice (Fig. 6e,f). Moreover, we found no significant difference in the frequency of dendritic Ca²⁺ spikes between SNI and sham mice (P = 0.63; sham daily CNO: 2.4 ± 0.35 events per active dendrite in 2.5 min; SNI daily CNO: 2.6 ± 0.39 events per active dendrite in 2.5 min). These results indicate that chronic activation of SOM cells in S1 with daily CNO treatment reduces the dendritic and somatic Ca²⁺ activity of L5 pyramidal neurons.

Notably, when we measured paw withdrawal threshold over time (Fig. 6a), we found that SOM cell activation for 1 week mitigated the mechanical allodynia in SNI mice as compared to saline-treated controls (Fig. 6g). One to 3 weeks after the last CNO injection at day 7, the pain threshold of SNI mice was not significantly different from that of sham mice, indicating that continuous activation of SOM cells with daily CNO treatment for 1 week prevents the formation of persistent pain in SNI mice.

DISCUSSION

While considerable efforts have been made towards unraveling the maladaptive changes in the spinal cord after peripheral nerve damage, much less is known about how cortical circuits are reorganized in chronic pain conditions. Using in vivo two-photon Ca²⁺ imaging, we found that pyramidal neurons located in L2/3 and L5 of S1 were hyperactive in a mouse model of neuropathic pain. We further found that increased synaptic inputs, dendritic Ca²⁺ spike generation, and changes of the activity of three major populations of cortical GABAergic interneurons collectively promoted pyramidal neuron hyperactivity (Supplementary Fig. 10). In addition, continuous activation of SOM-expressing interneurons suppressed the hyperactivity of pyramidal neurons and mechanical allodynia over 1 month. These findings indicate that dysfunctional inhibitory circuits contribute
Pyramidal neuron hyperactivity in neuropathic pain

Both L2/3 and L5 pyramidal neurons in S1 became profoundly activated after peripheral nerve injury (Fig. 1 and Supplementary Figs. 1 and 2). When considering ascending sensory information flow through S1, L2/3 neurons are classically positioned upstream of L5 neurons; information flows from thalamic inputs to L4 neurons, from L4 to L2/3 and then from L2/3 to L5. However, recently it has been shown that thalamic neurons can directly activate deep cortical layers. Therefore, both L2/3 and L5 neurons of S1 can be activated by ascending sensory information from the thalamus, and both layers eventually transmit information out of S1 to other brain areas associated with the processing of pain information, including subcortical regions for descending pain modulation. Future use of genetic approaches to manipulate neurons in specific cortical layers will aid in further delineating the contribution of L2/3 versus L5 neurons to the development of neuropathic pain.

One noteworthy finding of the current study is that the increased spontaneous activity of L5 pyramidal neurons in S1 is not reduced by acute silencing of peripheral sciatic nerves at 2 weeks after SN1 (Fig. 1f). These results suggest that peripheral nerve discharges are not the sole component driving abnormal cortical activity after the development of the chronic pain. Previous studies have demonstrated relief of spontaneous pain by systemic or spinal analgesia in animal models of neuropathic pain. Future study will be needed to better understand to what degree the maintenance of S1 hyperactivity in chronic pain condition requires inputs from the peripheral or spinal inputs.

Dendritic Ca²⁺ spike generation in neuropathic pain

By imaging apical tuft dendrites of L5 pyramidal neurons, we found a substantial and persistent increase in dendritic spine Ca²⁺ activity and NMDA-receptor-dependent dendritic Ca²⁺ spikes after peripheral nerve injury. Blockade of synaptic and dendritic activity in L1 of S1 reduced L5 pyramidal neuron hyperactivity. These results suggest that elevated synaptic inputs and the persistence of Ca²⁺ spike generation are likely to play a critical role in maintaining the hyperactive state of L5 cells and potentially L2/3 cells in neuropathic pain. It has been shown that the development of neuropathic pain is associated with long-term potentiation of synaptic transmission in ACC, a brain region mediating affective responses to painful stimuli. Dendritic Ca²⁺ spikes have been shown to induce long-lasting changes in synaptic plasticity (long-term potentiation and long-term depression) of existing synapses. It is thus possible that, following SN1, the generation of dendritic Ca²⁺ spikes may play a role in synaptic potentiation and lead to the increased Ca²⁺ activity of dendritic spines in S1. Dendritic Ca²⁺ spikes may also amplify synaptic inputs and increase the output of pyramidal neurons in SNI mice. Additionally, local Ca²⁺ spikes could potentially induce long-lasting changes by promoting formation of new dendritic spines, thus further contributing to persistent hyperactivity in S1 after peripheral nerve injury.

The mechanisms underlying the elevated generation of dendritic Ca²⁺ spikes in apical tuft dendrites remain to be determined. Our studies suggest that synaptic inputs to apical tuft dendrites are more synchronized in SNI mice as compared to sham mice. Such changes of synaptic input patterns favor dendritic Ca²⁺ spike generation. Notably, work in the ACC shows that peripheral nerve injury does not induce NMDA receptor-dependent dendritic Ca²⁺ spikes in the apical tuft of L5 pyramidal neurons, but rather is capable of generating fast local sodium dendritic spikes. L5 pyramidal neuron hyperexcitability in this brain region is linked to dysfunctional HCN (hyperpolarization-activated cyclic-nucleotide-gated) channels in apical tuft dendrites. It would be worthwhile to investigate how such changes of intrinsic excitability contribute to the generation of dendritic Ca²⁺ spikes and hyperactivity of L5 pyramidal neurons in S1. Moreover, as discussed below, reduced dendritic inhibition also promotes the generation of dendritic Ca²⁺ spikes and hyperactivity of L5 pyramidal neurons in S1.

Impaired cortical inhibition in neuropathic pain

Increasing evidence suggests that disinhibition of pain pathway is important to the development and maintenance of chronic pain. In the spinal cord, a loss of inhibitory neurotransmission has been implicated in several forms of chronic pain. Recent work in the ACC suggests that the loss of inhibitory synapses onto excitatory pyramidal neurons and the loss of the excitatory drive onto inhibitory fast-spiking interneurons creates a disinhibited cortical network. In S1, peripheral nerve injury causes an overall increase in the activity of GABAergic interneurons that is ineffective in dampening excitatory neuron activity due to changes in potassium–chloride cotransporter expression. However, it is unknown whether different types of GABAergic interneurons may exhibit different changes in activity following peripheral nerve injury. In this study, we used two-photon Ca²⁺ imaging to examine the activity of three groups of cortical interneurons in S1 of the awake mice. We found that peripheral nerve injury caused cell-type-specific changes in S1, in which VIP cells were activated and SOM and PV cell activity was downregulated. These changes developed within 1 week following peripheral nerve injury and persisted for more than 1 month. Previous studies have shown that three major subtypes of interneurons are connected in such a way that (i) VIP cells preferentially inhibit SOM cells, (ii) SOM cells strongly inhibit all other populations, and (iii) PV cells inhibit each other but poorly inhibit other populations. Our findings are consistent with these findings and suggest that the activity of inhibitory circuits is shifted to SOM/PV cells to increase excitatory activity in S1 during the development and maintenance of neuropathic pain (Supplementary Fig. 10). It would be worthwhile to test whether manipulating this inhibitory circuitry toward pyramidal cell hyperactivity would be sufficient to elicit pain behaviors in naive mice.

In contrast to the effect of manipulating SOM cells, we found that activation of PV cells was ineffective in altering the mechanical threshold for pain. This difference could be related to the facts that (i) PV cells primarily synapse perisomatically and have minimal regulation on dendritic Ca²⁺ spikes and (ii) PV cells may provide brief inhibition on pyramidal cell somata, whereas SOM cells exhibit high levels of spontaneous activity to inhibit apical dendrites of pyramidal neurons. It is important to note that besides for three major subtypes of interneurons in S1, other interneurons located in L1, which do not express SOM, PV or VIP markers, are known to target apical tuft dendrites, and their activation can suppress Ca²⁺ spike generation in L5 pyramidal neurons of S1. Future studies will be needed to examine whether L1 interneuron dysfunction is also involved in pyramidal neuron hyperactivity during the development of chronic pain.

Targeting inhibitory interneurons to modulate the development of chronic pain

Our study provides to our knowledge, the first direct evidence that impaired SOM cell activity is involved in the development of neuropathic pain. Given the reduction of SOM neuronal activity in SNI mice, we acutely manipulated the activity of S1 SOM cells and...
demonstrated that SOM cell activation was sufficient to decrease L5 dendritic and somatic Ca\(^{2+}\) hyperactivity. Notably, daily activation of SOM cells following SNi diminished L5 pyramidal neuron hyperactivity at the subcellular and cellular levels, as well as reducing mechanical allodynia. Hence, our findings suggest that manipulating interneuron activity after peripheral nerve injury could be an important avenue for the prevention of pyramidal neuron overexcitation and the transition from acute postoperative pain to chronic centralized pain.

Interneurons are becoming attractive targets for disease intervention via small molecule medications because of their profound impact on cortical circuit dynamics. For example, SOM cells express distinct receptors that have potent effects on neuronal network activity. Activation of SOM neuropeptide receptors has been proposed as one avenue for silencing neuronal networks in the face of epilepsy, which may be worth evaluating as a strategy to treat neuropathic pain states. Aside from medications, transcranial magnetic stimulation (TMS), a technique that uses high-intensity magnetic field and a brief electric current to create stimulation, was found to suppress apical tuft dendritic Ca\(^{2+}\) activity in L5 neurons. The use of TMS has been applied to many neurological diseases, such as schizophrenia, pain, depression and epilepsy, where interneuron dysfunction is thought to be a fundamental defect of the circuit. Whether or not TMS can be used to modulate activity of different interneuron classes (SOM or VIP cells) would be of great interest in developing new treatments for chronic pain.

**METHODS**

Methods, including statements of data availability and any associated accession codes and references, are available in the online version of the paper.

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**AUTHOR CONTRIBUTIONS**

J.C., T.J.B., W.-B.G. and G.Y. wrote the manuscript. J.C., T.J.B., W.-B.G. and G.Y. designed the experiments. J.C. and G.Y. performed the experiments, J.C. analyzed the data. All authors contributed to data interpretation. J.C., W.-B.G. and G.Y. wrote the manuscript.

**COMPETING FINANCIAL INTERESTS**

The authors declare no competing financial interests.

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ONLINE METHODS

Experimental animals. Transgenic mice expressing GCaMP6s in layer V (L5) pyramidal neurons, Thy1-GCaMP6 slow founder line 1, were generated at New York University School of Medicine. AAV experiments were conducted with C57BL/6 (Charles River Laboratory), Thy1-Cre (Jackson Laboratory; stock no. 006143), PV-IRES-Cre (Jackson Laboratory; stock no. 008069), SOM-IRES-Cre (Jackson Laboratory; stock no. 010344) and VIP-IRES-Cre mice (Jackson Laboratory; stock no. 010908). Mice were group-housed in temperature-controlled rooms on a 12-h light-dark cycle and were randomly assigned to different treatment groups. One to 3-month-old animals of both sexes were used for all the experiments. All mice were maintained at the New York University Old Public Health animal facility. Animal research was approved by Institutional Animal Care and Use Committee (IACUC).

Spared nerve injury and von Frey test. Spared nerve injury (SNI) of the sciatic nerve was performed on C57BL/6 adult (8–12 weeks) mice24. In brief, surgeries were performed under strict sterile conditions. Mice were deeply anesthetized and a small incision in the left thigh was made to expose the sciatic nerve and axotomy and ligation of the tibial and common peroneal nerves was performed, leaving the sural nerve intact. Great care was taken to avoid any contact with or stretching of the intact sural nerve. Muscle and skin were closed in two layers. For sham surgery, the sciatic nerve was exposed but not ligated or cut. The von Frey test was used to assess the onset and maintenance of mechanical allodynia over time. In all animal groups, mechanical allodynia threshold was examined using an electronic von Frey anesthesiometer (IITC Inc., Life Science Instruments) that measures the precise minimum pressure at which paw withdrawal occurs. Specifically, a von Frey tip of suitable rigidity was attached to an electronic probe and used to apply an increasing pressure to the lateral plantar aspect (the sural nerve skin territory) of the hind paw. The anesthesiometer displays the pressure at which the mouse retracts from the von Frey tip. Three trials for paw withdrawal were recorded for each day tested and an average was reported. The von Frey test was performed during the light cycle by the same researcher, who was blinded to the surgery, genotypes and treatments (CNO versus saline).

Surgical preparation for imaging awake, head-restrained mice. Dendritic and somatic imaging was carried out in awake, head-restrained mice in a quiet resting state27. Surgery preparation for awake animal imaging included attaching a head holder and creating a cranial window 51. Specifically, mice were deeply anesthetized with an intraperitoneal injection of ketamine (100 mg/kg) and xylazine (15 mg/kg) or isoflurane. The mouse head was shaved and the skull was exposed without damaging the temporal and occipital muscles. A head holder composed of two parallel micro-metal bars was attached to the animal’s skull with glue (Loctite 495) to help restrain the animal’s head and reduce motion-induced artifacts during imaging. A small skull region (~0.2 mm in diameter) located over the primary somatosensory cortex (S1) on the basis of stereotaxic coordinates (0.5 mm posterior and 1.5 mm lateral to the bregma) was removed and a round glass coverslip (approximately the same size as the bone being removed) was glued to the skull. Dental acrylic cement was applied to surrounding area to secure the metal bars, taking care not to cover the glass region with cement.

Upon awakening, mice with head mounts were habituated three times (10 min each time) in a custom-built imaging platform to minimize potential stress effects of head restraint and imaging. In this device, the animal’s head-mount was secured to metal blocks such that the head was fixed and perpendicular to the two-photon objective. The animal’s body was allowed to rest against the bottom of the solid imaging plate. There was no instrumentation to either of the animal’s hind limbs during imaging. Imaging experiments were started ~12–24 h after window implantation and free from anesthetic effects.

Ca2+ Imaging in dendrites and somata of L5 pyramidal neurons expressing GCaMP6s. The genetically encoded Ca2+ indicator GCaMP6 slow (GCaMP6s) was used for Ca2+ imaging of L5 pyramidal somata, apical dendrites, dendritic spines, and somata and axons of interneurons in S1. We used Thy1-Cre and Thy1-GCaMP6s (founder line 1) mice for pyramidal neuron imaging and PV, SOM, VIP Cre lines for interneuron imaging. For Cre mice, GCaMP6s was expressed with recombinant adeno-associated virus under the CAG promoter (AAV, serotype 2/1; ~10^12 GC/ml titer; produced by the University of Pennsylvania Gene Therapy Program Vector Core). To ensure better spread of AAV throughout S1, about 1–0.2 µl of AAV viruses were diluted 10x in artificial cerebrospinal fluid (ACSF) and injected (Picospritzer III; 15 p.s.l., 12 µm, 0.8 Hz) over ~10–15 min into L2/3 or 5 of S1 using a glass microelectrode around the coordinates of 0.5 mm posterior, 2 mm lateral to the bregma. In Thy1-GCaMP6s mice, the average total GCaMP6s signal during recordings was slightly lower than that in Thy1-Cre mice injected with AAV-GCaMP6s as a result of the relative amounts of GCaMP622.

Acute and chronic SOM cell activation was accomplished with Cre-dependent DREADD-hm3Dq under the human SYN1 promoter in SOM-Cre mice (AAV, serotype 2/1; ~10^12 GC/ml titer; produced by UNC Vector Core). For validation of SOM activation (DREADD-hm3Dq) or inactivation (DREADD-hm3Dq,Di in vivo (Fig. 5a,b,g), two viruses (AAV2/1-CAG-Flex-GCaMP6s and AAV2/1-HSYN-DIO-hmDq-mcherry or AAV2/1-HSYN-DIO-hmDq-Di-mcherry) were mixed at equal volumes and injected into S1 of SOM-ires-Cre mice, and SOM cell activity was imaged before and 20 min after CNO intraperitoneal injection. In experiments where SOM cells were activated and pyramidal cells were imaged (Figs. 5 and 6), AAV2/1-HSYN-DIO-hmDq-mcherry was injected into SOM-ires-Cre-Thy1-GCaMP6s mice. Similarly, in experiments where SOM cells were inactivated and pyramidal cells were imaged (Fig. 5i,j), AAV2/1-HSYN-DIO-hmDq-Di-mcherry was injected into SOM-ires-Cre-Thy1-GCaMP6s mice. As a control for DREADD expression, S1 was injected with AAV2/1-CMV-TurboFP (produced by the University of Pennsylvania Gene Therapy Program Vector Core) (Supplementary Fig. 8).

To evaluate the expression patterns of GCaMP6 and DREADD-Gq, SOM mice were fixed with 4% paraformaldehyde and the brain was sectioned at 200 µm thickness and immunostained for GFP (Abcam, ab39701:1:300) and RFP (Rockland, 600-401-3795, reactive against mCherry, 1:300) (Supplementary Figs. 6 and 9). Confocal microscopy (Zeiss LSM 700 confocal microscope; 10x air objective, numerical aperture 0.3) revealed that the average number of SOM cells in L2/3 of S1 infected with AAV2/1-HSYN-DIO-hmDq-mcherry was 205 ± 28 cells per 1 mm² (sham, 8 sections) and 194 ± 34 cells per 1 mm² (SNI, 9 sections) (Supplementary Fig. 6). The average number of SOM cells in L2/3 of S1 infected with AAV2/1-FLEX-CAG-GCaMP6s was 235 ± 21 cells per 1 mm² (sham, 12 sections) and 265 ± 21 cells per 1 mm² (SNI, 15 sections). PV mice infected with AAV2/1-HSYN-DIO-hmD3Dq-mcherry in S1 had a labeling density of 110 ± 24 cells per 1 mm² (sham, 6 sections) and 149 ± 13 cells per 1 mm² (SNI, 7 sections) (Supplementary Fig. 9). SOM mice infected with AAV2/1-CMV-TurboFP showed expression restricted to S1. Dimensions of the infected area (Supplementary Fig. 8) were similar between SNI and sham mice. Transgenic mice infected with AAV were prepared for head fixation and imaging after 2 weeks of AAV expression. In vivo two-photon imaging was performed with an Olympus Fluoview 1000 two-photon system equipped with a Ti:sapphire laser (MaiTai DeepSee, Spectra Physics) tuned to 920 nm. The average laser power on the sample was ~20–30 mW for imaging in the L1 of the cortex. All experiments were performed using a 25x objective immersed in an ACSF solution and with a 2x (somata) and 4x (dendrites) digital zoom. All images were acquired at frame rates of 2–10 Hz (2 µs pixel dwell time). Image acquisition was performed using FV10-ASW v2.0 software and analyzed post hoc using NIH Image software.

Data analysis. During quiet resting, motion-related artifacts (derived from the animal’s respiration and heart beat) were typically less than 2 µm as detected in our cortical measurements. Vertical movements were infrequent and minimized by habituation, the use of two micro-metal bars attached to the animal’s skull (described above) by dental acrylic, and the use of a custom-built body platform. If the animal struggled in the body platform, imaging time points from those segments were excluded from quantification. All imaging stacks were registered using NIH ImageJ plug-in StackReg.

L2/3 and L5 somata. In this study, we used GCaMP6s, an indirect reporter of neuronal spiking activity53. All cells labeled with GCaMP6s expressed GCaMP6 in the cytoplasm, and cells that had nuclear expression were excluded from analysis. Regions of interests (ROIs) corresponding to visually identifiable somata (pyramidal cells and interneurons) were selected for quantification at different cortical depths from the pial surface. Imaging planes were from L2/3 and L5, which corresponded to cells positioned 200–300 µm from pial surface and cells

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more than 600 µm from the pial surface, respectively. Cells were not recorded from layer 4 in this study. The fluorescence time course in each cell body was measured with NIH ImageJ by averaging all pixels within the ROI covering the soma. The ΔF/F₀ was calculated as ΔF/F₀ = (F − F₀)/F₀ × 100, where F₀ is the baseline fluorescence signal averaged over a 2-s period corresponding to the lowest fluorescence signal over the 2.5-min recording period.

Comparing calcium traces to neuronal firing rates. GCaMP6s can produce large fluorescence transients (~20% ΔF/F₀) even in response to single action potentials, and individual spikes within a burst result in stepwise fluorescence increases53. However, when neuronal firing rates are high, it becomes difficult to resolve the number of action potentials owing to the long decay time constant of GCaMP6 fluorescence. We found that there was a diversity in calcium traces of pyramidal cells in SNI and sham mice, which likely reflects burst and nonburst firing of L5 neurons54,55. Although it is difficult to report firing rates on the basis of calcium responses within individual cells, several studies using two-photon imaging and patch-clamp electrophysiology have found a significant correlation between calcium signals and action potential generation in L2/3 neurons54,55. In vivo, dendritic Ca²⁺ spikes were expressed using the peak fluorescence transients (~20% ΔF/F₀) even in response to single action potentials, and individual spikes within a burst result in stepwise fluorescence increases53. However, when neuronal firing rates are high, it becomes difficult to resolve the number of action potentials owing to the long decay time constant of GCaMP6 fluorescence.

Dendritic spines. In Figure 2, dendritic spine Ca²⁺ transients were defined as those with changes of fluorescence (ΔF/F₀) >35% for GCaMP6s during 1-min imaging session. The threshold was more than 3 times the s.d. of baseline fluorescence noise for GCaMP6s. Spine coactivity was assessed by applying Pearson correlation coefficients to the fluorescence signal of the spine head across the 1-min recordings from random spine pairs on dendritic segments.

Drugs. Local application of drugs. MK801 (M107, Sigma-Aldrich) and TTX citrate (1069, Tocris) were applied to the surface of S1 after removing a small bone flap (~200 µm in diameter) adjacent to a thinned skull window. MK801 (100 µM) and TTX (100 nM) were dissolved directly in ACSF to final concentrations. The bone flap for drug delivery was made during awake head mounting and covered with a silicone elastomer such that it could be easily removed at the time of imaging. Because small molecules diffuse rapidly in the cortex, we estimated that the drug concentration was reduced ~10-fold in the imaged cortical region, such that the final effective concentration would be ~10 µM and 10 nM for MK801 and TTX, respectively. As a control, we applied ACSF after removing a bone flap. In Figure 1f, TTX (100 nM) was locally delivered to the surgical site of sciatic nerve.

In Supplementary Figure 1, MK801 (15 µg/ml in saline; 0.1 or 0.25 µg/g body weight) was delivered by intraperitoneal injection.

DREADD activation of SOM cells. DREADD activation was achieved in SOM-IRESCre mice in which AAV2/1-HSYN-DIO-hM4Di-mCherry was injected into S1 2 weeks before the Ca²⁺ imaging. AAV-CMV-TurboRFP was used as a control AAV vector. Clozapine N-oxide (CNO, C0832, Sigma-Aldrich) was dissolved in saline to a concentration of 0.5 mg/ml. In Figure 5, Ca²⁺ imaging of dendritic tufts and somata was performed during a quiet resting state without CNO. Twenty minutes after CNO was administrated by intraperitoneal injection to each mouse (0.3 ml per 30 g body weight), the same cortical region was reimaged for second session (another 2.5 min recording). Mechanical threshold was also assessed before and 20 min after CNO injection. In Figure 6, CNO or saline was injected daily over the first week (a total of seven injections) following SNI. Consecutive CNO injections were spaced 24 h apart and mechanical threshold was measured 1 h before each CNO injection (or 23 h after previous CNO injection). No more CNO injections were given after the first week. Ca²⁺ imaging of dendritic tufts and somata was performed at day 8.

Statistics. Summary data are presented as means ± s.e.m. No statistical methods were used to predetermine sample sizes, but our sample sizes are similar to those reported in previous publications27. Data distribution was assumed to be normal, but this was not formally tested. The variance was similar between groups that were being statistically compared. Tests for differences between two populations were performed using unpaired or paired t test. Two-way ANOVA followed by Bonferroni’s or Tukey’s test was also used to compare significance between various groups. Significant levels were set at P ≤ 0.05. All statistical analyses were performed using GraphPad Prism. A Supplementary Methods Checklist is available.

Data availability. The data supporting the findings of this study are available from the corresponding author upon reasonable request.