Burst and Tonic Spinal Cord Stimulation Both Activate Spinal GABAergic Mechanisms to Attenuate Pain in a Rat Model of Chronic Neuropathic Pain

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Abstract

Background: Experimental and clinical studies have shown that tonic spinal cord stimulation (SCS) releases gamma-aminobutyric acid (GABA) in the spinal dorsal horn. Recently, it was suggested that burst SCS does not act via spinal GABAergic mechanisms. Therefore, we studied spinal GABA release during burst and tonic SCS, both anatomically and pharmacologically, in a well-established chronic neuropathic pain model.

Methods: Animals underwent partial sciatic nerve ligation (PSNL). Quantitative immunohistochemical (IHC) analysis of intracellular GABA levels in the lumbar L4 to L6 dorsal spinal cord was performed after 60 minutes of burst, tonic, or sham SCS in rats that had undergone PSNL (n = 16). In a second pharmacological experiment, the effects of intrathecal administration of the GABAA antagonist bicuculline (5 μg) and the GABAβ antagonist phaclofen (5 μg) were assessed. Paw withdrawal thresholds to von Frey filaments of rats that had undergone PSNL (n = 20) were tested during 60 minutes of burst and tonic SCS 30 minutes after intrathecal administration of the drugs.

Results: Quantitative IHC analysis of GABA immunoreactivity in spinal dorsal horn sections of animals that had received burst SCS (n = 5) showed significantly lower intracellular GABA levels when compared to sham SCS sections (n = 4; P = 0.0201) and tonic SCS sections (n = 7; P = 0.0077). Intrathecal application of the GABAA antagonist bicuculline (5 μg; n = 10) or the GABAβ antagonist phaclofen (5 μg; n = 10) resulted in ablation of the analgesic effect for both burst SCS and tonic SCS.

Conclusions: In conclusion, our anatomical and pharmacological data demonstrate that, in this well-established chronic neuropathic animal model, the analgesic effects of both burst SCS and tonic SCS are mediated via spinal GABAergic mechanisms.

Key Words: gamma-aminobutyric acid, burst spinal cord stimulation, GABAβ receptor antagonist, peripheral nerve injury, mechanical hypersensitivity
INTRODUCTION

Tonic spinal cord stimulation (SCS) is a last-resort treatment method for patients that suffer from intractable chronic neuropathic pain.\textsuperscript{1–6} The standard tonic SCS protocol consists of continuous, tonic, electrical stimulation applied to the dorsal columns of the spinal cord with a frequency within the range of 40 to 80 Hz and a pulse width between 200 and 500 \(\mu\)S.\textsuperscript{2,7,8} The concept of SCS emerged as a direct application of the Gate Control Theory of Melzack and Wall in 1965.\textsuperscript{7,9} It was postulated that antidromic stimulation of the non-nociceptive \(A_\beta\) fibers could close the “spinal gate,” located in the dorsal horn of the spinal cord. Closing of the “gate” is facilitated by inhibitory interneurons located in the superficial laminae of the dorsal horn and it is believed that the neurotransmitter gamma-aminobutyric acid (GABA) plays a pivotal role in this process.\textsuperscript{8,10,11} During the development of neuropathic pain, Janssen and colleagues observed, in a partial sciatic nerve ligation (PSNL) model, increased intracellular levels of GABA in the dorsal horn.\textsuperscript{12} Later, it was demonstrated that tonic SCS decreased intracellular GABA immunoreactivity (IR) in the dorsal horn of these rats that had undergone PSNL.\textsuperscript{13} At the same time, extensive experimental microdialysis-work has demonstrated that tonic SCS increases extracellular GABA levels in the dorsal horn of allodynic rats that had undergone PSNL.\textsuperscript{14–16} Thus, GABAergic interplay seems to be an important aspect of the analgesic mechanisms underlying tonic SCS in the PSNL model of chronic neuropathic pain. The role of GABA in segmental SCS mechanisms was further elucidated by the administration of pharmacological agents that specifically modulate GABA release in the dorsal horn during tonic SCS in neuropathic rats that had undergone PSNL.\textsuperscript{14–16} Local perfusion with a GABA\(_B\) receptor antagonist in the dorsal horn transiently abolished the SCS-induced effect in neuropathic rats,\textsuperscript{16} and rats not receiving adequate pain relief with tonic SCS (nonresponders) were turned into responders by administration of the GABA\(_B\) receptor agonist baclofen.\textsuperscript{17} The aforementioned preclinical findings were successfully translated to the clinic, where patients with neuropathic pain who had experienced a deficient tonic SCS effect had improved pain relief following the intrathecal administration of baclofen,\textsuperscript{18} further confirming the theory that local spinal GABAergic mechanisms are pivotal for the effects of tonic SCS. While GABA plays a role in the underlying mechanism of tonic SCS, the neurotransmitters involved in other stimulation paradigms, such as burst SCS, have not been clearly identified. Because patients show different responses to different stimulation paradigms, the underlying mechanisms may be different.\textsuperscript{19} The burst waveform consists of closely spaced pulses delivered in a packet or burst, directly followed by a quiescent period or interburst interval.\textsuperscript{20} Burst SCS has proven to be effective in patients with failed back surgery syndrome and radiculopathy, and clinical trials have demonstrated its ability to help patients to reduce their analgesic intake.\textsuperscript{21–23} In addition, burst SCS reduces neuropathic pain without generating paresthesia in the affected limb or area.\textsuperscript{24–26} Yet, from a mechanistic point of view, the burst waveform is still in its infancy. Results of source-localized electroencephalographic studies and patient questionnaires suggest that burst SCS preferentially activates the medial pain pathway, and hence modulates emotional-affective pain aspects.\textsuperscript{25,27} On a segmental level, an experimental electrophysiological study has aimed to elucidate the involvement of GABA in the spinal mechanism underlying burst SCS.\textsuperscript{28} It was found that the presence of a GABA\(_B\) receptor antagonist blocked attenuation of dorsal horn neuronal firing during tonic SCS but not burst SCS. Furthermore, blood serum GABA measurements showed that systemic GABA levels were not increased following burst SCS. From this, the researchers concluded that burst SCS might not act via spinal GABAergic mechanisms.\textsuperscript{29} However, it should be mentioned that the aforementioned study was performed not only in an uncommon rat model for chronic neuropathic pain, the painful cervical nerve root compression model, but these experiments were also terminal; thus, no behavioral testing during the conscious state of the animals was performed.\textsuperscript{29} This makes interpretation of these data in the context of understanding the role of spinal GABA in a chronic neuropathic pain model difficult, as most experimental data on pain relief and the spinal GABAergic mechanism underlying tonic SCS have been documented and studied in the well-described and validated PSNL model or a similar nerve injury model for peripheral mononeuropathy.\textsuperscript{13–17,30} Therefore, in order to further understand the spinal mechanism underlying burst SCS we aimed to study, both anatomically and pharmacologically, the role of GABA in behavior and pain-relieving mechanisms underlying burst and tonic SCS, in a well-established chronic neuropathic PSNL model.
METHODS

Ethics Statement

The experiments were performed in accordance with the European Directive for the Protection of Vertebrate Animals Used for Experimental and Other Scientific Purposes (86/609/EU). The protocol was approved by the Animal Research Committee of the Maastricht University Medical Centre (Project License number 2017-022).

Animals

All experiments were performed using male Sprague Dawley rats ($n=36$), which were young adults (5 weeks of age) at the start of the experiment (150 to 200 g). Sixteen animals were used in the first experiment, in which we assessed the quantitative immunohistochemical (IHC) analysis of GABA IR in spinal dorsal horn sections of animals after SCS ($n=16$). Twenty animals were used in the second experiment, in which we intrathecally administered GABA$_{A/B}$ antagonists before SCS in order to assess the effects of these pharmacological agents on the behavioral pain-relieving effects of SCS ($n=20$). Animals were housed in groups of 2, in polycarbonate cages in a climate-controlled vivarium maintained under controlled temperature (21°C±1°C), relative humidity (55°C±15%), and artificial lighting (12:12 reversed light/dark cycle) with distilled water and rodent food available ad libitum. The vivarium was equipped with a central radio system, continuously producing background music at 45 decibels, in order to desensitize the animals for experimenter-related noise. All procedures were conducted between 09:00 and 16:00 hours. Experiments were conducted during the dark, active phase of the rodent circadian rhythm.

Partial Sciatic Nerve Ligation

A unilateral ligation of the left sciatic nerve was performed as described by Seltzer et al.,$^{30}$ and previously applied in our laboratory.$^{29,31–33}$ In short, animals were anesthetized with 3% to 5% isoflurane (Abbott Laboratories Ltd., Kent, U.K.) and air enriched with 100% oxygen at a constant flow rate of 250 mL/min. By use of an automatic heating pad, body temperature was maintained at 37.5°C. The nervus ischiadicus from the left hind paw was exposed by blunt dissection and carefully freed from surrounding connective tissue. Subsequently, the nerve was partially ligated (by approximately one third) using 8/0 non-absorbable silk sutures. The wound was then closed with 4/0 silk sutures. Development of mechanical hypersensitivity (mechanical allodynia) was monitored with the use of von Frey assays for 14 consecutive days. At day 14, the presence of mechanical hypersensitivity was confirmed if the log (50% paw withdrawal threshold [PWT]) was decreased by 0.2 units compared to baseline (day 0).$^{34}$

Assessment of Mechanical Hypersensitivity (von Frey Assay)

Mechanical hypersensitivity was assessed by use of von Frey filaments. Von Frey assessment was always conducted in the same room, isolated from external sounds and equipped with artificial red light sources (temperature: 21 ± 1°C; relative humidity: 55 ± 15%). Prior to testing, animals were placed in the behavioral set-up for 15 minutes in order to acclimate to the new surroundings. The assessment room was equipped with a mobile radio, continuously producing background music at 45 decibels. PWTs to von Frey filaments were assessed using the up-down method.$^{35}$ Von Frey filaments of linearly incrementing stiffness (bending forces 0.6, 1.2, 2.0, 3.6, 5.5, 11.1, 8.5, 15.1 and 28.84 g) were applied to the plantar surface of the hind paw of the rats for 5 consecutive seconds. A negative response (the hind paw was not withdrawn) was followed by the subsequent filament with greater bending force. In contrast, a positive response (the hind paw was withdrawn) was followed by application of the previous filament with a lower bending force. After completion of a sequence of 6 consecutive responses, the 50% PWT was calculated.$^{35}$ The predetermined cut-off value was set at 28.84 g. For statistical analysis, the 50% PWTs were logarithmically transformed to yield a linear scale.

Tissue Preparation

For immunohistochemistry experiments, 16 animals in experiment 1 were divided into 3 groups: sham SCS ($n=4$), tonic SCS ($n=7$), and burst SCS ($n=5$). Animals were killed 60 minutes after the start of (sham) SCS. Tissue perfusion was performed transcardially with a mixture of 4% paraformaldehyde and 15% picric acid in 0.2 M phosphate-buffered saline (PBS; pH 7.6) after anesthesia with pentobarbital (100 mg/kg body weight). Then, lumbar spinal cord regions L4 to L6 were removed by a laminectomy, post-fixed overnight at 4°C, and cryoprotected for 24 hours in 10% sucrose. This was followed by at least 72 hours’ incubation in

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25% sucrose (in 0.1 M PBS; pH 7.6) at 4°C. Subsequently, tissues were frozen in solid carbon dioxide. Thirty-micrometer thick transverse cryosections were mounted on gelatine-coated glass slides and stored at −20°C until staining procedures were performed.

**Immunohistochemical Detection of GABA**

The immunohistochemistry protocol was performed as described by Janssen et al. 13 In short, staining procedures were performed at room temperature unless stated otherwise. First, glass slides were air dried for 2 hours and subsequently washed in Tris-buffered saline (TBS, 0.1 M, pH 7.6) including 0.3% Triton X-100 (TBS-T). Blocking was performed with 2% normal donkey serum (Sigma-Aldrich, Zwijndrecht, The Netherlands). The serum was diluted in TBS-T and applied for a 1-hour incubation period. Then, the sections were incubated with a polyclonal rabbit anti-GABA antibody (1:5,000 diluted in TBS-T; Sigma-Aldrich, A2052) for a time period of 48 hours. After 48 hours, the excess primary antibody was removed by use of TBS, after which sections were incubated with the secondary alexa fluor 488 donkey anti-rabbit IgG antibody (1:100 diluted in TBS-T; Invitrogen, Breda, The Netherlands, A21206) for 2 hours. Subsequently, sections were rinsed with TBS and coverslipped with TBS/glycerol (20%/80%).

**Quantification of Immunostaining**

Quantitative IHC analysis of spinal dorsal horn GABA staining was performed as described by Janssen et al. 12,13 Photomicrographs were taken of both ipsilateral and contralateral lumbar L4 to L6 spinal cord immunostained sections using a Provis AX70 fluorescent microscope (Olympus, Hamburg, Germany). The microscope was connected to a digital black and white video camera (U-CMAD-2; Olympus), equipped with CellP software. Rexed laminae 1 to 3 of the dorsal horn were determined as regions of interest for the GABA IR analysis. 36 Grayscale values were calculated for these laminae. Analysis of gray scale spinal cord pictures was performed by a blinded observer using the AnalySIS software program CellP (Soft Imaging Systems, Münster, Germany).

**Implantation of Spinal Cord Stimulation Device**

Implantation of the SCS device was performed according to the standard protocol used in our institution. 29,32,33,37,38 In short, a small laminectomy was made at T13, after which the spinal cord was exposed by use of a surgical rotary tool. A custom-made cylindrical 4-contact lead (0.72 mm diameter; Boston Scientific Neuromodulation, Valencia, CA, U.S.A.) was inserted into the epidural space in the caudal direction. Electrode configuration was set at alternating cathode and anode settings (rostral to caudal: + - + -). Then, the electrode was secured to a spinous process with tissue adhesive (Histoacryl; B Braun Medical BV, Oss, The Netherlands) to prevent electrode migration. The electrode wires were tunneled subcutaneously to the neck of the animal, and the stimulator connectors were attached with 4/0 silk sutures. Animals were given 2 days for recovery prior to the initiation of SCS experiments.

**Intrathecal Implantation**

After successful implantation of the SCS device, 20 animals in experiment 2 underwent a small laminectomy at T12, after which the spinal cord was exposed by use of a surgical rotary tool. The membrane was carefully opened, after which a polyethylene catheter from Instech Laboratories (Plymouth Meeting, PA, U.S.A.; 27 gauge, length 1.5 cm) was inserted and tunneled intrathecally to the level of the L3 to L5 spinal segments as described by Truin et al. 31 The catheter was secured to a spinous process with tissue adhesive (Histoacryl; B Braun Medical BV) to prevent migration. Subsequently, the catheter was slowly flushed with 10 μL of saline and the wound was closed with 4/0 silk sutures. Animals with signs of paralysis directly after surgery were excluded. Correct placement of the catheter was confirmed when lidocaine 2% injection (10 mg/mL) resulted in paralysis or dragging of the hind limbs. 31

**Spinal Cord Stimulation**

For stimulation of the dorsal columns, an A-M Systems stimulator (MultiStim: Programmable 8-Channel Stimulator, model 3800, 220 V/50 Hz) fitted with a stimulus isolator (model 3820 for A-M Systems MultiStim) was used. The stimulator was set to deliver constant current biphasic stimulation for both the tonic and burst SCS modes. The tonic SCS motor threshold (MT) was determined at the following settings: pulse width of 200 μS administered at a frequency of 2 Hz. Burst SCS MT was determined at the following settings: pulse width of 1,000 μS, 5 pulses (449 Hz intraburst frequency) administered at an interburst frequency of
2 Hz. The amplitude was gradually increased until symmetrical contractions of the hind limbs were perceived by hand and/or visually observed. Then, either a tonic SCS paradigm at 66% of MT (frequency 50 Hz, pulse width 200 μS), or a biphasic burst SCS paradigm at 50% MT (interburst frequency 40 Hz, pulse width 1,000 μS, 5 spikes at 449 Hz intraburst frequency) was applied for 60 minutes. The SCS parameters were based on previously determined optimal settings for pain relief with burst SCS (50% MT) and tonic SCS (66% MT). In all experiments, the PWTs to von Frey filaments were assessed before the start of SCS treatment (baseline), at 15, 30, 45, and 60 minutes after stimulation was turned on. The investigator was blinded to the stimulation condition throughout the whole experiment.

**Intrathecal Administration of GABA<sub>ABA</sub>B Receptor Antagonists (Experiment 2)**

The concentration of the GABA<sub>ABA</sub>B receptor antagonists bicuculline (5 μg; ≥97%; Sigma-Aldrich) and phaclofen (5 μg; ≥97%; Sigma-Aldrich) was chosen based on literature demonstrating a dose-response curve for both antagonists. Because the peak dorsal horn drug concentrations occurred 30 minutes after intrathecal administration, antagonists were applied to the spinal cord 30 minutes before SCS and von Frey testing protocols were initiated. Intrathecal administration of the GABA<sub>ABA</sub>B receptor antagonists (5 μg) was followed by the administration of 20 μL saline. Vehicle administration consisted of 10 μL saline followed by the administration of 20 μL saline.

**Data Analysis**

Paw withdrawal threshold's to von Frey filaments are presented as mean ± standard error of the mean (SEM). In line with previous Von Frey studies, von Frey data were logarithmically transformed to obtain a linear scale and to account for Weber’s Law. For statistical analysis of differences in the withdrawal thresholds over time within the groups of experiments 1 and 2, the nonparametric Friedman test was used, followed by Dunn’s post hoc test. For the analysis of differences in the withdrawal thresholds between groups (ipsilateral and contralateral withdrawal thresholds), the nonparametric Mann–Whitney U-test was used. Before statistical analysis of GABA IR gray values in experiment 1, a Shapiro-Wilk normality test was performed. Then, the Kruskal-Wallis test was performed in order to assess statistical differences over group means. Dunn’s Multiple Comparison Test followed this in order to assess significant differences between the group means. All statistical analyses were performed with α = 0.05 using IBM SPSS statistics version 23 (IBM Corp., Armonk, NY, U.S.A.).

**RESULTS**

**Development of Mechanical Hypersensitivity (von Frey) and SCS**

In experiment 1, after PSNL surgery, ipsilateral hindpaw PWTs were significantly lower than contralateral hindpaw PWTs ($P = 0.019$); all 16 animals qualified as hypersensitive due to increased response to mechanical stimulation by von Frey filaments (contralateral average PWTs: 11.2 ± 0.6 g (day post lesion [DPL] 7) vs. 1.1 ± 0.5 g [DPL 7]; $P = 0.0049$) (see Methods) (Figure 1A). Burst SCS ($n = 5$) significantly increased PWTs compared to baseline PWTs at 15 ($P = 0.046$), 30 ($P = 0.025$), 45 ($P = 0.016$), and 60 minutes of SCS ($P = 0.019$) (Figure 2A). For tonic SCS ($n = 7$), PWTs significantly differed from baseline PWTs at 15 ($P = 0.027$), 30 ($P = 0.026$), 45 ($P = 0.033$), and 60 minutes of SCS ($P = 0.031$) (Figure 2A). Sham SCS ($n = 4$) did not increase PWTs at any time points (see Figure 2A).

In experiment 2, after PSNL surgery, ipsilateral hindpaw PWTs were significantly lower than contralateral hindpaw PWTs ($P = 0.0081$) at DPL 7; all 20 animals qualified as hypersensitive (ipsilateral average PWTs: 10.9 ± 1.1 g [pre-lesion] vs. 1.3 ± 0.7 g [post-lesion]; $P = 0.0064$) (see Methods) (Figure 1B). For burst SCS ($n = 10$), PWTs significantly decreased from baseline PWTs at 15 ($P = 0.041$), 30 ($P = 0.027$), 45 ($P = 0.028$), and 60 minutes of SCS ($P = 0.022$) (Figure 2B). For tonic SCS ($n = 10$), PWTs significantly decreased from baseline PWTs at 15 ($P = 0.029$), 30 ($P = 0.028$), 45 ($P = 0.031$), and 60 minutes of SCS ($P = 0.035$) (see Figure 2B). No significant differences were reported between development of mechanical hypersensitivity in experiments 1 and 2. In addition, no significant differences were reported between SCS time points in experiments 1 and 2. When the SCS data of experiments 1 and 2 were pooled, burst SCS ($n = 15$) and tonic SCS ($n = 17$) significantly increased PWTs compared to baseline PWTs at 15 ($P = 0.042$ and $P = 0.028$, respectively), 30 ($P = 0.027$ and $P = 0.026$, respectively), 45 ($P = 0.021$ and $P = 0.032$, respectively).
respectively), and 60 minutes of stimulation ($P = 0.025$ and $P = 0.033$, respectively) (Figure 2C). Burst SCS PWTs and tonic SCS PWTs significantly differed at 15 minutes of SCS ($P = 0.038$) (see Figure 2C).

In experiment 1, anti-GABA IHC revealed an intense IR predominantly in laminae 1 to 3 of the lumbar spinal dorsal horn with clear identification of GABA IR cell bodies (Figure 3B). Gray scale values were calculated for these laminae. The Shapiro-Wilk normality test was not passed ($P = 0.0395$). Accordingly, a Kruskal-Wallis test was performed, which showed a significant difference between mean gray values for the sham SCS, tonic SCS, and burst SCS groups ($P = 0.0085$, Kruskal-Wallis statistic = 9.524). Statistical testing of GABA staining intensity in the spinal dorsal horn and gray values revealed that the tonic SCS group, although it showed a strong tendency, did not differ significantly from the sham SCS group ($P = 0.1609$). The burst SCS group differed significantly from the sham SCS group ($P = 0.0201$) and from the tonic SCS group ($P = 0.0077$).

In experiment 2 (Figure 4), thresholds for mechanical hypersensitivity were not affected after 15 minutes of tonic SCS with intrathecal administration of the GABA $\alpha$ antagonist bicuculline, as PWTs were significantly increased as compared to baseline ($P = 0.044$) (Figure 5A). Meanwhile, at 30 minutes ($P = 0.38$), 45 minutes ($P = 0.65$), and 60 minutes ($P = 0.32$) of tonic SCS, thresholds for mechanical hypersensitivity were affected, since PWTs did not significantly differ from baseline PWTs. Administration of the GABA $\alpha$ antagonist bicuculline affected thresholds for mechanical hypersensitivity at all time points during burst SCS.
PWTs did not significantly differ from baseline PWTs at 15 minutes ($P = 0.74$), 30 minutes ($P = 0.49$), 45 minutes ($P = 0.52$), and 60 minutes of burst SCS ($P = 0.67$) (see Figure 5A).

After intrathecal administration of the GABA$_B$ antagonist phaclofen, thresholds for mechanical hypersensitivity did not differ from baseline for both tonic SCS and burst SCS (Figure 5B). PWTs at 15 minutes ($P = 0.35$ and $P = 0.38$, respectively), 30 minutes ($P = 0.56$ and $P = 0.41$, respectively), 45 minutes ($P = 0.43$ and $P = 0.50$, respectively), and 60 minutes of SCS ($P = 0.64$ and $P = 0.59$, respectively) did not significantly differ from baseline PWTs (see Figure 5B).

In a control experiment, administration of vehicle (saline minus antagonist) was administered. Thresholds for mechanical hypersensitivity were not affected, since PWTs significantly increased over time for both tonic SCS ($n = 10$) and burst SCS ($n = 10$) at 15 minutes ($P = 0.01$ and $P = 0.01$, respectively), 30 minutes ($P = 0.01$ and $P = 0.01$, respectively), 45 minutes ($P = 0.01$ and $P = 0.03$, respectively), and 60 minutes ($P = 0.01$ and $P = 0.01$, respectively) as compared to baseline (Figure 5C).

Administration of the GABA$_A$ antagonist bicuculline ($n = 5$) and the GABA$_B$ antagonist phaclofen ($n = 5$) 30 minutes prior to sham SCS did not increase PWTs as compared to baseline PWTs (Figure 5D).

Administration of the GABA$_A$ antagonist bicuculline ($n = 5$) and the GABA$_B$ antagonist phaclofen ($n = 5$) 30 minutes prior to Von Frey testing decreased PWTs of the contralateral paw (unaffected by the sciatic nerve ligation) at 15 minutes ($P = 0.01$ and $P = 0.01$, respectively), 30 minutes ($P = 0.01$ and $P = 0.01$, respectively), 45 minutes ($P = 0.01$ and $P = 0.03$, respectively), and 60 minutes ($P = 0.01$ and $P = 0.01$, respectively) of Von Frey testing as compared to PWTs 30 minutes before baseline (Figure 6).
**DISCUSSION**

IHC analysis of dorsal horn sections showed a decrease in intracellular GABA levels immediately after burst SCS. Intracellular GABA levels were significantly decreased compared to sham SCS animals and tonic SCS animals. In a second experiment, administration of a GABA\textsubscript{A} and GABA\textsubscript{B} receptor antagonists abolished the analgesic effect of both tonic SCS and burst SCS. Based on these findings, we conclude that the analgesic

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**Figure 4.** Timeline of experiment 2: pharmacological intrathecal experiments. SCS, spinal cord stimulation.

| Day 1                      | Day 2                      | Day 3                      | Day 4                      |
|---------------------------|----------------------------|----------------------------|----------------------------|
| -No Anta + Tonic SCS (n=5)| -Tonic + GABA\textsubscript{A}-anta (n=3) | -Vehicle + Tonic (n=5)     |
| -No Anta + Burst SCS (n=5)| -Burst + GABA\textsubscript{A}-anta (n=3) |                             |
| -GABA\textsubscript{A}-anta + Sham SCS (n=5) | -Tonic + GABA\textsubscript{A}-anta (n=2) |                             |
| -GABA\textsubscript{B}-anta + Sham SCS (n=5) | -Burst + GABA\textsubscript{A}-anta (n=2) |                             |

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**Figure 5.**

A. The effects of tonic spinal cord stimulation (SCS) (n = 10) and burst SCS (n = 10) on paw withdrawal thresholds (PWTs) based on sensitivity to von Frey filaments 30 minutes after GABA\textsubscript{A} antagonist bicuculline administration. PWTs were assessed at baseline and at 15, 30, 45, and 60 minutes of SCS. B. The effects of tonic SCS (n = 10) and burst SCS (n = 10) on PWTs 30 minutes after GABA\textsubscript{A} antagonist phaclofen administration. PWTs were assessed at baseline and at 15, 30, 45, and 60 minutes of SCS. C. The effects of tonic SCS (n = 5) and burst SCS (n = 5) on PWTs 30 minutes after vehicle administration. PWTs were assessed at baseline and at 15, 30, 45, and 60 minutes of SCS. D. The effects of sham SCS and GABA antagonist bicuculline (n = 10) or phaclofen (n = 10) administration on PWTs 30 minutes after antagonist administration. The dotted line represents the average PWT baseline prior to sciatic nerve ligation.

\*P < 0.05 for burst SCS vs. baseline; \*P < 0.05 for tonic SCS vs. baseline.
effect of burst SCS is mediated via a spinal GABAergic mechanism. GABAergic interneurons are known to be important players in the pain gate system that is located in the dorsal horn of the spinal cord,\textsuperscript{39,42,43} which in its turn is a fundamental component of the working mechanisms underlying tonic SCS.\textsuperscript{8–11,13–17,44} Both experimental and clinical studies have documented the importance of segmental GABA release by turning SCS nonresponders into responders via intrathecal administration of the GABA\textsubscript{B} agonist baclofen (in subeffective doses).\textsuperscript{16–18} The majority of experimental studies on the effects of tonic SCS were performed in sciatic nerve injury models, including the sciatic nerve ligation model (PSNL) and the chronic constriction injury model.\textsuperscript{13,16,17,29,31–33,37,38,45} Therefore, in order to adequately compare and correlate our findings to previous literature, we deliberately chose to perform our experiments in the partial sciatic nerve ligation model.\textsuperscript{13} Initially, we found that intracellular GABA levels in the rat dorsal horn were significantly decreased following 60 minutes of burst SCS, hence indicating that burst SCS does induce the release of GABA in the dorsal horn. Our notion was further confirmed by the results of our second experiment, where we demonstrated that the GABA\textsubscript{A} antagonist phaclofen and GABA\textsubscript{B} antagonist bicuculline both abolished the pain-relieving behavioral effect of burst and tonic SCS. Our findings on the effects and mechanisms of tonic SCS are in line with earlier work performed at our laboratory, where it was demonstrated that tonic SCS decreased intracellular GABA IR in the dorsal horn of rats that had undergone PSNL.\textsuperscript{13} Perhaps even more important is that, in line with our study, experimental pharmacological work has previously shown that local perfusion with a GABA\textsubscript{B} receptor antagonist in the dorsal horn of neuropathic rats transiently abolished the tonic SCS-induced effect.\textsuperscript{16} Meanwhile, preclinical studies conducted in a sciatic nerve injury model demonstrated that intrathecal administration of the GABA\textsubscript{B} agonist baclofen and the GABA\textsubscript{A} agonist muscimol, administered intrathecally, resulted in a marked and long-lasting increase of withdrawal thresholds in rats not achieving adequate pain relief with tonic SCS (nonresponders), although muscimol produced a less prominent threshold increase as compared to baclofen.\textsuperscript{46} This and our data show that tonic SCS operates by potentiating the spinal GABAergic systems in general, and that the tonic SCS effect depends on both GABA\textsubscript{A} receptor and GABA\textsubscript{B} receptor signaling, although the SCS effect seems to be more linked to the GABA\textsubscript{B} receptor system.\textsuperscript{45} In our study, administration of a GABA\textsubscript{A} and GABA\textsubscript{B} receptor antagonist prevented the pain-relieving effect of burst SCS.

**Figure 6.** Paw withdrawal thresholds (PWTs) based on sensitivity to von Frey filaments of the contralateral paw 30 and 15 minutes before bicuculline (n = 10) or phaclofen administration (n = 10), and at baseline and 15, 30, 45, and 60 minutes after antagonist administration. The dotted line represents the average PWT baseline prior to sciatic nerve ligation (*p < 0.05 for bicuculline vs. –30 minutes; †p < 0.05 for phaclofen vs. –30 minutes). SCS, spinal cord stimulation; WT, withdrawal threshold.
SCS. Our conclusion that the pain-relieving effect of burst SCS is mediated via spinal GABAergic mechanisms may conflict with findings reported by Crosby and colleagues. Based on electrophysiological analysis of neuronal firing after burst SCS, these investigators reported that the presence of a GABA \(_B\) receptor antagonist did not block the attenuation of dorsal horn neuronal firing, when SCS was applied at 90% of the MT. Based on these findings the authors conclude that burst SCS does not act via a spinal GABAergic mechanism, whereas tonic SCS does. However, because the electrophysiological experiments were terminal, no behavioral testing was performed during the conscious state of the animals. Yet, behavioral analysis and thus assessment of the pain-relieving effect is known to be the most important indicator of a treatment or compound’s translational value. Furthermore, even though the experiments provide novel insights into the working mechanisms underlying tonic and burst SCS, the fact that the experiments were performed in a painful cervical nerve root compression rat model makes it difficult to compare the findings to the majority of experimental SCS literature, since most experimental data on pain relief and spinal GABAergic mechanisms underlying tonic and burst SCS have been documented and studied in peripheral nerve injury models for mononeuropathy. It is interesting that in the study by Crosby and colleagues, both burst and tonic SCS was applied at 90% MT, an intensity known to induce unwanted side effects in the animals. Our study and analysis was based on previously determined optimal settings for pain relief with burst SCS (50% MT) and tonic SCS (66% MT). Moreover, according to the strength-duration curve, with SCS administered at 90%, MT is above the perception threshold, while burst SCS in the clinical setting is usually applied below the perception threshold (paresthesia free). Experimental evidence and clinical observations indicate that burst SCS has a delayed wash-in effect as compared to tonic SCS. An experimental study by Meuwissen et al. demonstrated that after 15 minutes, burst SCS does induce pain relief, but at a suboptimal level as compared to the plateau phase, which is reached after 30 minutes (when burst and tonic SCS reach similar levels of pain relief). It has been proposed that burst SCS is capable of modulating both the medial and the lateral pain pathways. As stated by De Ridder, “the exact mechanism of the selective routing is unknown but could be related to diameter differences at a bifurcation, in which only the burst waveform is powerful enough to overcome the higher transmission resistance of the smaller C-fibers that make up the medial pain pathway.” Meanwhile, evidence shows that tonic SCS, via descending pathways, exerts an inhibitory GABAergic effect in the spinal cord. Therefore, it could be hypothesized that 5 minutes of burst SCS, as applied in the Crosby et al. study, was not sufficient to set in motion the delayed supraspinal mechanisms that lead to optimal pain relief as observed after the effects of burst SCS have plateaued (30+ minutes). Additionally, after only 5 minutes of burst SCS, Crosby and colleagues might have observed a GABA-independent segmental mechanism of burst SCS, unaffected by the GABA \(_B\) antagonist, while the supraspinal loop did not yet set in motion the descending pathways that exert an inhibitory effect via GABAergic mechanisms in the spinal cord. It should be mentioned that in our study we applied biphasic active recharge burst SCS, while Crosby and colleagues applied passive recharge burst SCS. With monophasic SCS, the pulse is given in only one direction from one electrode to the other. With a biphasic pulse, direction of the pulse is reversed by changing the polarity of the electrodes, allowing the system to actively recharge after each pulse. There are indications that biphasic stimulation is more safe; however, whether or not there are physiological or clinical differences between the biphasic and monophasic burst waveforms is unknown. To date, no data from clinical or preclinical studies have been performed that have directly compared passive recharge burst SCS and active recharge burst SCS; therefore, no conclusive statements regarding the efficacy of these burst waveforms can be made. However, future research that aims to elucidate the working mechanisms of, and (possible) differences between, these waveforms is desired. Interestingly, our results demonstrate that blocking of the GABA \(_A\) receptor does not prevent a significant increase in PWTs after 15 minutes of tonic SCS, while the effect was successfully suppressed during the remaining minutes of stimulation. This suggests that tonic SCS possesses the ability to, initially, activate GABA-independent pain-relieving mechanisms. It is believed that the supraspinal mechanisms of tonic SCS arise from activation of the fast-conducting lateral pain pathway. Once activated, supraspinal areas activated by the lateral pain pathway may modulate via descending pathways the pain signals coming in on a segmental level, through, for example, the release of serotonin and norepinephrine at the spinal dorsal horns.
CONCLUSION

In conclusion, our data, based on experiments in a validated commonly used experimental chronic neuropathic pain model, demonstrate that spinal GABA plays a role in the working mechanisms of burst SCS in the attenuation of pain. Further analysis of the GABAergic mechanisms underlying burst SCS at different time points could shed more light on the exact mechanisms at play.

CONFLICTS OF INTEREST

Jianwen Wendy Gu and Tianhe C. Zhang are employees of Boston Scientific Inc., and Elbert A.J. Joosten is a consultant for Boston Scientific Inc. Koen Meuwissen and Luuk de Vries have no conflicts of interest to declare.

FUNDING

This work was supported by a sponsored research contract (ISRNO060009) from Boston Scientific Inc., granted to Prof. Dr. E.A.J. Joosten.

AUTHOR CONTRIBUTIONS

K.M. performed the experiments, analyzed the data, and wrote the manuscript. L.d.V. processed post-mortem tissue and analyzed gray value data. J.W.G., T.C.Z., and E.J. edited the manuscript and contributed to the design of the experiment.

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