Long-Term Spinal Cord Stimulation Alleviates Mechanical Hypersensitivity and Increases Peripheral Cutaneous Blood Perfusion in Experimental Painful Diabetic Polyneuropathy

Maarten van Beek, PhD; Denise Hermes, BSc; Wiel M. Honig, BSc; Bengt Linderoth, PhD; Sander M. J. van Kuijk, PhD; Maarten van Kleef, PhD; Elbert A. Joosten, PhD

Objectives: This study utilizes a model of long-term spinal cord stimulation (SCS) in experimental painful diabetic polyneuropathy (PDPN) to investigate the behavioral response during and after four weeks of SCS (12 hours/day). Second, we investigated the effect of long-term SCS on peripheral cutaneous blood perfusion in experimental PDPN.

Methods: Mechanical sensitivity was assessed in streptozotocin induced diabetic rats (n = 50) with von Frey analysis. Hypersensitive rats (n = 24) were implanted with an internal SCS battery, coupled to an SCS electrode covering spinal levels L2–L5. The effects of four weeks of daily conventional SCS for 12 hours (n = 12) or Sham SCS (n = 12) were evaluated with von Frey assessment, and laser Doppler imaging (LDI).

Results: Average paw withdrawal thresholds (PWT) increased during long-term SCS in the SCS group, in contrast to a decrease in the Sham group (Sham vs. SCS; p = 0.029). Twenty-four hours after long-term SCS average PWT remained higher in the SCS group. Furthermore, the SCS group showed a higher cutaneous blood perfusion during long-term SCS compared to the Sham group (Sham vs. SCS; p = 0.048). Forty-eight hours after long-term SCS, no differences in skin perfusion were observed.

Discussion: We demonstrated that long-term SCS results in decreased baseline mechanical hypersensitivity and results in increased peripheral blood perfusion during stimulation in a rat model of PDPN. Together, these findings indicate that long-term SCS results in modulation of the physiological circuitry related to the nociceptive system in addition to symptomatic treatment of painful symptoms.

Keywords: Experimental research, laser Doppler imaging, long-term spinal cord stimulation, mechanical hypersensitivity, painful diabetic polyneuropathy, physiology, spinal cord stimulation, vasodilation, von Frey

Conflict of Interest: The authors reported no conflict of interest.

INTRODUCTION

Spinal cord stimulation (SCS) is a successful last resort treatment modality for patients suffering from various chronic pain disorders. Currently, SCS is mainly prescribed as a symptomatic treatment of painful symptoms. It can result in successful pain relief in patients with chronic pain arising from various disorders, including painful diabetic polyneuropathy (PDPN) (1–5). Besides symptomatic treatment, there are numerous indications that continuous neuremodulation of the spinal cord results in physiological changes (6–11). These physiological changes are key players in understanding the working mechanism behind SCS and identifying targets for improvement. This is of importance since many patients do not respond to SCS treatment or fail to respond over time, despite the technological advances that have been made in the last decades (12–14). SCS mechanisms can be investigated clinically to some extent; however important aspects need a more detailed insight at the physiological and molecular level which can only be obtained from animal experiments. However, there are several limitations in the translation of...
experimental studies to the clinical situation. Probably one of the most important limitations is the relatively short duration of stimulation (short-term SCS) that is applied in experimental models. Although these short-term SCS paradigms have proven to be useful in the assessment of pain related behavior and investigating the acute physiological changes in the nociceptive network (7,15–19), the need for insight into long-term (clinically more relevant) SCS effects and related physiological changes is urgent.

Recent developments in SCS technology have resulted in the production of smaller pulse generators, which allows continuous stimulation of the spinal cord in rats for hours, days, weeks, or even months (20,21). For instance, long-term SCS (6 hours/day for 3 months) in a model of spared nerve injury (SNI) resulted in decreased mechanical hypersensitivity and increased activity levels (20). Moreover, mechanical hypersensitivity remained decreased during long-term SCS throughout the study, whereas the initially increased activity levels later returned toward values as observed in rats receiving sham stimulation. No cumulative effect or loss of effect of long-term SCS on mechanical hypersensitivity was observed. Considering the vascular mechanisms that are involved in PDPN pathology, the effect of long-term SCS on cutaneous blood perfusion is an important aspect to investigate. Short-term experimental SCS has been shown to result in very local improvements in cutaneous blood perfusion in diabetic rats (6). Clinical improvements in patients with ischemic pain after long-term SCS suggest that long-term SCS might result in improved blood perfusion of a larger area in the affected limb (11,22). Besides repetitive short-term SCS in experimental PDPN (23), long-term SCS has not yet been established for experimental PDPN.

The first aim of this study was to investigate pain behavior, based on paw withdrawal thresholds (PWT) to von Frey filaments, during and after four weeks of daily SCS (12 hours/day). Second, we investigated the effect of long-term SCS on cutaneous blood perfusion in the hind limbs using laser Doppler imaging (LDI).

METHODS

Animals

All experiments were performed using male Sprague-Dawley rats (n = 60), which were eight weeks of age at the start of the experiment (300–350 g). Animals were housed on a reversed day night rhythm in transparent plastic cages with access to food and water ad libitum. The experiments were approved by the Animal Research Committee of the Maastricht University Medical Centre.

Induction of Diabetes Mellitus

Diabetes mellitus (DM) was induced in Week 2 with a single intraperitoneal (i.p.) injection of 65 mg/kg of streptozotocin (STZ) (Sigma-Aldrich, Schnelldorf, Germany) (n = 50). Control rats (n = 10) were injected with saline. Before STZ injection, rats were weighed and fasted overnight. STZ was freshly dissolved in sterile 0.9% NaCl to a solution of 65 mg/mL. Four days after STZ injection, blood glucose level was determined in blood derived from the saphenous vein using a standard blood glucose meter (Accu-Chek Aviva, Roche Diagnostics GmbH, Mannheim, Germany). Rats with a glucose level of ≥15 mmol/L were considered diabetic (24). Throughout the study additional blood glucose measurements were regularly performed to confirm the status of DM. When glucose levels exceeded 30 mmol/L, we placed a third of a slow releasing insulin pellet (Lin-Shin Canada, Inc.) subcutaneously in the trunk only in the first two weeks after induction of DM in order to decrease general somatic dysfunction caused by severe DM during the first phase of the study.

Implantation of Spinal Cord Stimulation Electrode

Diabetic rats with a decrease in PWT of ≥0.2 unit in log(50%PWT) as compared with baseline were selected for implantation of an internal SCS device (PDPN rats). A change of 0.2 unit in log(50%PWT) corresponds to a 50% chance of rats responding to the next higher or lower filament (25,26). Rats that did not show a decrease of ≥0.2 unit in log(50%PWT) were excluded from the study. The implantation of the quadripolar SCS electrode was performed according to previous work (23) and adapted to the use of an Implantable Pulse Generator (IPG). In short, under general anesthesia, a small laminectomy was made at vertebrate level L1 via laminectomy and placed over L2–L5 spinal levels.
Behavioral Assessment

Mechanical hypersensitivity was assessed according to the “up-down method” (25). In short, von Frey filaments with incrementing stiffness (bending forces 1.2, 2.0, 3.6, 5.5, 8.5, 15.1, and 28.84 g) were applied to both hind paws of the rats for 5 sec through the wire mesh floor of the examination cage. If the hind paw was not withdrawn (=negative response), the next filament with higher bending force was applied, whereas the previous filament with lower bending force was applied if the hind paw was withdrawn (positive response). The average 50% PWT (50%PWT) was calculated after completion of a sequence of six consecutive responses. A cut-off value of 28.84 g was defined. Calculated 50%PWTs were logarithmically transformed to a linear scale for statistical analysis. PWTs were assessed before the induction of PDPN, before implantation of the SCS device, and weekly after implantation. Twenty-four hours after cessation of long-term SCS, PWTs to von Frey filaments were assessed before, during, and after an acute stimulation paradigm of 40 min (Fig. 2). The experimenter was blinded during all behavioral analyses and rats were tested in random order.

Statistical Analyses

Longitudinal differences between groups in bodyweight, blood glucose levels, MTs, and withdrawal thresholds to Von Frey filaments before and during long-term SCS were analyzed using a linear mixed effects regression model with a random intercept for each rat. The random intercept ensures that the repeated measures within each rat are correctly accounted for. In addition, the linear mixed effects model analyzes all rats, despite missing data due to removal of the SCS device or electrode malfunction over time. Because of complete data, differences in withdrawal thresholds to Von Frey filaments in Week 11 were analyzed using a Repeated Measures Analysis of Variance (RM-ANOVA). Similar to the linear mixed effects model the RM-ANOVA accounts for repeated measures within each rat. We defined time (five levels: 0, 15, 30, 60, and 90 min) as the with-subjects factor and group allocation (i.e., Sham, SCS, or control) was assigned as between-subjects factor. Differences in cutaneous blood perfusion during long-term stimulation were analyzed by means of Analysis of Variance (ANOVA). All analyses were performed with the Statistical Package for the Social Sciences, version 23 (SPSS Inc., Chicago, IL, USA). p-Values of ≤0.05 were considered to indicate statistical significance.

RESULTS

Induction of Diabetes Mellitus

Intraperitoneal injection of 65 mg/kg STZ (n = 50) resulted in blood glucose levels ≥15 mmol/L in 48 rats (96%). Twenty-four rats (48%) developed PDPN and were implanted with an internal SCS device. Blood glucose levels increased from 8.3 mmol/L (95% confidence interval [CI]: 7.9, 8.7) at baseline to 27.8 mmol/L (95% CI: 24.7, 30.9) and 26.7 mmol/L (95% CI: 24, 29.5) postinjection in the Sham (n = 12) and SCS group (n = 12), respectively. Blood glucose levels remained elevated throughout the rest of the study period (Fig. 3).

Bodyweight of control animals (n = 10) increased from 322 g (95% CI: 311, 333) at the start of the study to 485 g (95% CI: 458, 511) at this temperature has been shown to fall in the range of a thermoneutral zone in several rodent strains (28). Rats were given 10 min of acclimatization before the start of the first scan. Three scans (resolution x: 227, y: 50; speed: 4 msec/pixel, 1.35 min per scan) were performed per rat and the average flux units from both hind paws of the three scans were recorded and analyzed (Software: Moor LDI V5.3). LDI was performed at baseline, pre-SCS and in Week 10, and 11. The LDI measurement after long-term SCS in Week 11 was performed no more than 48 hours after cessation of stimulation (Fig. 2). Rats were scanned in random order.
the end of the study (Week 11). The SCS (n = 12) and Sham group (n = 12) underwent a decrease in bodyweight after STZ injection followed by an increase to 392 g (95% CI: 365, 420) and 372 g (95% CI: 352, 391), respectively, in Week 11 (Fig. 4). Linear mixed regression analysis revealed no significant differences in blood glucose levels or bodyweight between the Sham and SCS group.

Spinal Cord Stimulation

One rat in the Sham group was implanted with an IPG with battery failure and two more rats in the Sham group failed to show MTs in Week 10, due to removal of sutures by the rats and subsequent subcutaneous scratching of the silicon connection wire of the SCS device. Furthermore, one rat in the Sham group underwent considerable discomfort due to anterolateral migration of the IPG, which was therefore explanted in Week 11. Hence, nine and eight rats were included in the Sham group in Week 10 and 11, respectively, during all procedures requiring active electrical stimulation in the Sham group (i.e., determination of MT and the analysis of PWTs to von Frey filaments in Week 11). All rats in the SCS group (n = 12) successfully completed the study with a functional SCS system.

MT was determined in all PDPN rats with an implanted functional SCS device (SCS group (n = 12) and Sham group (n = 11) total n = 23). The amplitude (in Volts) required to induce a motor response in the hind limbs or lower trunk during anesthesia in the surgical procedure of SCS implantation was significantly higher compared with follow-up measurements in awake rats (p < 0.001).

During implantation, the average MT of all implanted rats was 1.40V (95% CI: 1.03, 1.78) during surgery, compared with 0.23V (95% CI: 0.21, 0.26) in Week 7, and 0.34V (95% CI: 0.28, 0.41) in Week 11. No statistical significant differences were observed between the Sham and SCS group (Fig. 5).

Behavioral Assessment

Average 50%PWT to Von Frey filaments before the induction of DM was 14.8 g (95% CI: 12.6, 17.4). Linear mixed effects regression analysis revealed a significant interaction between time and group during long-term SCS (p = 0.044), indicating that the effect of time differed between the group strata. The average 50%PWT in the SCS group increased from 4.8 g (95% CI: 3.1, 7.6) pre-long-term SCS to 9.8 g (95% CI: 5.9, 15.8) in Week 10, whereas average 50%PWT in the Sham group decreased from 7.1 g (95% CI: 5.1, 10.0) to 5.6 g (95% CI: 3.5, 8.9) (Sham vs. SCS; p = 0.029) (Fig. 6). The average 50%PWT in control rats remained stable over time throughout the study.

Repeated Measures Analysis of Variance revealed a significant overall effect of time during 40 min of SCS in both the SCS and Sham group (p < 0.001). During stimulation, PWTs peaked at 30 min with an average increase of 0.19 unit in log(50%PWT). Although no significant differences between Sham and SCS rats were noted, an absolute difference of 0.2 unit in log(50%PWT) was present at baseline (0 min). Average 50%PWT at baseline was 7.1 g (95% CI: 4.8, 10.5) in the SCS group, whereas average 50%PWT in the Sham group was 4.5 g (95% CI: 2.7, 7.1). Average 50%PWT at 90 min was 6.1 g (95% CI: 4.9, 7.5) in the SCS group and 4.9 g (95% CI: 3.2, 7.5) (Fig. 7).

Cutaneous Blood Perfusion

Both the SCS and Sham group showed lower blood perfusion of the plantar surface of the hind paws after induction of DM and before long-term SCS (p = 0.028). The SCS group showed a significantly higher blood perfusion of the hind limbs as compared with the Sham group at the end of four weeks long-term SCS (p = 0.048). SCS rats showed an increase from 60.6% of baseline blood perfusion (95% CI: 44.6, 76.5) before long-term SCS (pre-SCS) to 70.5% of baseline blood perfusion (95% CI: 58.1, 83.0) in Week 10. In contrast, Sham rats showed a minimal decrease from 53.0% of baseline blood perfusion (95% CI: 39.7, 66.4) to 52.6% of baseline blood perfusion.
After long-term SCS no differences were observed between the SCS and Sham group (SCS: 47.3\% of baseline blood perfusion [95\% CI: 32.7, 61.9] and Sham: 44.5\% of baseline blood perfusion [95\% CI: 33.2, 55.8]) (Fig. 8). Control rats did not show significant changes in blood perfusion during the study (pre-SCS: 98.5\% of baseline blood perfusion [95\% CI: 69.1, 127.8] and in Week 10: 102.1\% of baseline blood perfusion [95\% CI: 71.0, 133.1]).

**DISCUSSION**

The first aim of the present study was to examine the effect of long-term SCS (four weeks daily, 12 hours/day) on pain behavior in experimental PDPN. Second, we investigated the effect of long-term SCS on cutaneous blood perfusion in the hind paws of PDPN rats.

To study the effect of long-term SCS on pain behavior in experimental PDPN we performed von Frey assessments before, during, and after long-term SCS. Analysis of these data revealed a statistically significant interaction between time and group during long-term SCS. Mean absolute PWT increased 0.31 unit log(50\%PWT) in the SCS group during four weeks of long-term SCS, whereas the average PWT in the Sham group decreased 0.08 unit log(50\%PWT). This is of importance, since a difference of 0.2 unit log(50\%PWT) can be considered as a relevant difference in sensory perception (25,26), and served as inclusion criterion for SCS implantation in our previous experiments (23,27). Twenty-four hours after cessation of long-term SCS, the SCS group shows a 0.2 unit log(50\%PWT) higher baseline PWT as compared with the Sham group, while no significant difference in PWT was observed between the groups when 40 min of SCS was applied. PWTs did decrease further after cessation of 40 min SCS in rats that received Sham long-term SCS compared with rats that received long-term SCS. These findings indicate that mechanical hypersensitivity is alleviated 24 hours after four weeks of 12 hours/day SCS, partially due to another reason than active SCS at that time point. Possibly, functional or even structural changes have occurred in the nociceptive system due to long-term SCS, which contribute to the alleviation of baseline mechanical hypersensitivity. Possible central mechanisms include inhibition of windup and prolonged synaptic depression in lamina II dorsal horn neurons to nociceptive inputs (29). In contrast, clinical reports usually report less SCS treatment success over time, which can ultimately result in explantation of the SCS device (4,5,30). A possible explanation for the discrepancy between these clinical reports and our experimental findings is that human patients with chronic pain might experience an adaptation phenomenon (4,31). Whereas pain relief in rats is scored by the application of a calibrated filament with an absolute force, patients are aware of the fact that they are receiving treatment and score their pain (on visual or numeric scales) based on previous conditions. Our data indicates that baseline withdrawal thresholds were higher after long-term SCS, and that acute short-term SCS did not significantly increase baseline PWTs further in rats treated with long-term SCS. It could therefore be speculated that the higher baseline PWTs in PDPN rats are in line with a patients’ baseline condition at a certain time point. The absence of further pain relief upon active stimulation might result in patients perceiving no treatment success, although they have an improved baseline condition.

As a second aim, we analyzed cutaneous blood perfusion data before, during, and after long-term SCS. Previous rat studies have demonstrated very local improvements of blood perfusion upon short-term SCS paradigms (8,32–34), which we could demonstrate with LDI as well. However, attempts to measure increased blood perfusion of the entire plantar surface of the hind paw failed during short-term SCS in both diabetic as well as healthy rats (unpublished results). Since after four weeks, a higher blood perfusion of the entire plantar surface of the hind paws can be demonstrated only

**Figure 6.** Effect of long-term SCS (12 hours/day from Week 7 to Week 10) on paw withdrawal thresholds (50\% PWT(grams) and log(50\%PWT)) as assessed with von Frey filaments. Mean ± SEM (a). Scatterplot of PWTs in the SCS and Sham group before long-term SCS and in Week 10. The horizontal lines mark the averages in each group (b). *p < 0.05 regression analysis.

**Figure 7.** Effect of long-term SCS and paw withdrawal thresholds (50\% PWT(grams) and log(50\%PWT)) as assessed with von Frey filaments in Week 11: A traditional 40 min stimulation paradigm was conducted in both the SHAM and SCS group 24 hours after cessation of long-term SCS. Mean ± SEM. *p < 0.01 rm-ANOVA.
during active stimulation, we could speculate that the total amount of stimulation explains the alleviation of baseline mechanical hypersensitivity, since the total duration of stimulation in this study is 336 hours. As long-term SCS results in clinical improvement and pain relief in patients with ischemic pain (11,22,35), a prolonged improved perfusion of the entire plantar surface of the rat hind paws might result in clinical improvement and subsequent pain relief as well. The baseline PWT to Von Frey filaments is presumably still affected 24 hours after cessation of long-term SCS, while no increased blood perfusion was present at that time. Given the fact that peripheral blood perfusion after cessation of long-term SCS did not differ between the Sham and SCS group, we conclude that long-term SCS had no effect on baseline blood perfusion. Instead, long-term SCS resulted in an increased potency of SCS to induce vasodilation. The increased blood perfusion of the hind paws during SCS is indicative for increased CGRP signaling, which occurs mainly via antidromic activation of small unmyelinated CGRPergic fibers (6,7,34,36,37). These findings might indicate that long-term SCS results in preservation or sprouting of CGRPergic small diameter fibers. Indeed, experimental studies support the hypothesis that electrical stimulation of nerve tissue results in axon sprouting, possibly via BDNF signaling (38–40). Moreover, Intra Epidermal Nerve Fiber Density and CGRP labeling has been shown to be affected in both clinical and experimental diabetes (41,42). If long-term SCS indeed results in improved or preserved functioning of small diameter nerve fibers, it might be worthwhile to initiate SCS therapy earlier instead of waiting until all other therapeutic efforts have failed. Indeed, clinical evidence underlines the importance of neuropathy stage for SCS treatment success in PDPN patients (3). Therefore, the effect of long-term SCS on CGRPergic nerve fibers in dermatomes that are targeted by SCS should be the future focus in experimental SCS studies in PDPN. Postmortem analysis of nervous tissue in the hind paws could confirm whether or not long-term SCS indeed results in the preservation of nervous tissue in dermatomes targeted by SCS. Furthermore, other vasodilatory pathways of SCS should be taken into account, such as sympathetic inhibition (43–45).

Out of 24 rats with an implanted SCS system, only one rat underwent severe discomfort related to implantation of the pulse generator. No infections occurred although STZ induced diabetes is accompanied by severe pathology, including increased susceptibility to infection (46–48). This might be explained by the fact that we provided short-term glycemic control, which can reduce surgical site infections (49). However, insulin was only administered in the two weeks after STZ injection to preserve DM status during the experimental phase of long-term SCS. Therefore, it is more likely that extensive surgical training prior to experimental surgeries, and critical evaluation of aseptic conditions during surgery, are responsible for the absence of infections in our PDPN rats. No further complications occurred as a result of implantation of the internal IPG. Based on these results, we conclude that implantation of an internal IPG is a suitable approach to study the effects of long-term SCS in experimental PDPN. Average MT was significantly higher in sedated rats during surgery compared with awake rats during long-term SCS (Fig. 5), which can be partially explained by the use of anesthesia as has been previously demonstrated (50). Furthermore, an increase of 0.11V in average MT was observed during the four weeks postsurgery. The thickness of the cerebrospinal fluid layer between the dura mater and spinal cord, which is anatomically different in rats in a prone position during surgery, is an important factor that contributes to electrical conductivity to Aβ fibers in the dorsal column and therefore determines the required SCS amplitude (51). Post surgery, MT was determined in awake rats, which might have led to an underestimation of MT during experimental procedures requiring anesthesia, i.e., determination of peripheral blood perfusion and nerve conductance velocity. The amplitude of SCS, which is a percentage of MT, has been shown to be positively correlated to local peripheral vasodilation (8), suggesting that the effect size of our blood perfusion data might be underestimated. Furthermore, small sample sizes in experimental studies limit the extrapolation of results to clinical trials including larger heterogeneous patient populations. To provide more insight of experimental long-term SCS on nociception in PDPN rats, a larger battery of pain assessment techniques could have been included in this study. However, the health status of PDPN rats is relatively poor and should be taken into account when designing long-term experiments including multiple assessment techniques, especially when sedation is involved.

In conclusion, our results indicate that long-term SCS, with conventional stimulation parameters, results in decreased mechanical
hypersensitivity even after cessation of SCS in experimental PDPN. Furthermore, blood perfusion of the hind limbs is significantly increased during long-term SCS, although this effect diminishes after cessation of SCS. Therefore we suggest that structural changes have occurred in the nociceptive network during long-term SCS in experimental PDPN and the effect of SCS on vascular reactivity might be causally involved.

Authorship Statements

Maarten van Beek was responsible for the execution of experiments, collection, analysis, interpretation, and documentation of the data. Denise Hermes and Wiel M. Honig assisted in executing experiments. Sander M. J. van Kuijk contributed to the analysis of the data. Bengt Linderoth, Maarten van Beek, and Elbert A. Joosten contributed to interpretation of the data and reviewed the manuscript.

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COMMENTS

van Beek, et. al, have performed an interesting study of the long-term effect of conventional SCS in a rat model of diabetic neuropathy. Such studies are important to investigate mechanisms of pain alleviation and will help to guide advances in SCS technology. The persistence of decreased mechanical hypersensitivity even after cessation of SCS brings up an important topic of neuroplasticity engendered by SCS. The authors have included both sham and control arms and should be commended on the quality of their investigation. As technology for SCS continues to evolve, greater research on the mechanisms behind pain reduction by SCS will help to expand indications and perhaps elucidate reasons for why different waveforms have different clinical efficacy.

Nestor Tomycz, MD
Pittsburgh, PA, USA

Applying long-term SCS which better mimics that used in the clinic is quite challenging in rodent models. The study is well designed and nicely executed, and hence the potential translational value of this study is high.

Yun Guan, MD
Baltimore, MD, USA

This is the first publication of long-term stimulation applied to animal models. The study clearly demonstrates that long-term SCS results in decreased baseline mechanical hypersensitivity and results in increased peripheral blood perfusion during stimulation in a rat model of PDPN. The study advances further our ability to mimic the clinical therapy in animal model.

Sam Eldabe, MD
Middlesbrough, UK

Comments not included in the Early View version of this paper.