REVIEW

463

Current Perspectives on Neurostimulation for the Management of Chronic Low Back Pain: A Narrative Review

This article was published in the following Dove Press journal: Journal of Pain Research

David A Provenzano (1)^{1,2} Jared A Heller (1)³ Michael C Hanes⁴

¹Pain Diagnostics and Interventional Care, Sewickley, PA, USA; ²Western PA Surgery Center, Wexford, PA, USA; ³Washington & Jefferson College, Washington, PA, USA; ⁴Jax Spine & Pain Centers, Jacksonville, FL, USA

Correspondence: David A Provenzano Pain Diagnostics and Interventional Care, 301 Ohio River Blvd Suite 203, Sewickley, PA, 15143, USA Tel +1 412-221-7640 Fax +1 412-490-7640 Email davidprovenzano@hotmail.com



Abstract: Neurostimulation techniques for the treatment of chronic low back pain (LBP) have been rapidly evolving; however, questions remain as to which modalities provide the most efficacious and durable treatment for intractable axial symptoms. Modalities of spinal cord stimulation, such as traditional low-frequency paresthesia based, high-density or high dose (HD), burst, 10-kHz high-frequency therapy, closed-loop, and differential target multiplexed, have been limitedly studied to determine their efficacy for the treatment of axial LBP. In addition, stimulation methods that target regions other than the spinal cord, such as medial branch nerve stimulation of the multifidus muscles and the dorsal root ganglion may also be viable treatment options. Here, current scientific evidence behind neurostimulation techniques have been reviewed with a focus on the management of chronic axial LBP.

Keywords: spinal cord stimulation, chronic, axial, neuromodulation

Introduction

Approximately one in five adults suffer from chronic low back pain (LBP) with over 575 million individuals suffering worldwide.^{1–3} Chronic LBP is associated with significant direct health-care costs including physical therapy, inpatient services, prescription medications, and interventional care, as well as indirect costs including lost work productivity.⁴

When discussing chronic LBP. it is important to provide a contextual definition. In reference to time, chronic back pain is typically defined as pain lasting longer than 12 weeks or beyond the expected period of healing.⁵ A well-defined underlying pathological cause may not always be identified. As defined by NASS (North American Spine Society), chronic LBP is "pain of musculoskeletal origin extending from the lowest rib to the gluteal fold that may at times extend as somatic referred pain into the thigh."⁶ Chronic pain that is localized in the lower back is considered axial LBP, while pain that radiates past the buttock and down the leg(s) is classified as radicular pain. Pain that chronically occurs concurrently in the back and the leg is considered back and leg pain. Further classifications can be subdivided into individuals that have chronic LBP following back surgery (eg postlaminectomy pain syndrome; PLPS) and those that have nonsurgical refractory LBP.

Journal of Pain Research 2021:14 463-479

© 2021 Provenzano et al. This work is published and licensed by Dove Medical Press Limited. The full terms of this license are available at https://www.dovepress.com/ terms.php and incorporate the Creative Commons Attribution — Non Commercial (unported, v3.0) License (http://creativecommons.org/license/by-nc/3.0/). By accessing the work you hereby accept the Terms. Non-commercial uses of the work are permitted without any further permission from Dove Medical Press Limited, provided the work is properly attributed. For permission for commercial use of this work, please see paragraphs 4.2 and 5 of our Terms (https://www.dovepress.com/terms.php).

Unfortunately, the treatment of chronic LBP can be challenging and refractory to many interventions. Forms of neuromodulation, such as spinal cord stimulation (SCS), have been utilized over the last 50 years to assist in the treatment of intractable LBP often with a radicular component. Since the first SCS unit was implemented in a clinical setting by Shealy, SCS has been growing in popularity as its clinical safety, cost utility, and efficacy increase.⁷⁻⁹ Spinal cord stimulation is now a widely accepted form of therapy for chronic intractable neuropathic pain and is the most commonly employed neurostimulation therapy for the treatment of chronic pain. More specifically, SCS has been used extensively for the treatment of PLPS (approximately 70% of all implants); however, it has not yet been fully investigated as a treatment modality for patients who have not previously undergone back surgery.^{10,11} When treating PLPS, SCS has traditionally been used to target radicular symptoms, but at times lacks effective prolonged coverage of axial LBP symptoms.^{11,12}

The use of neurostimulation for the treatment of chronic LBP has been met with mixed results. Due to a multitude of recent technological and hardware advancements, including programming (eg novel waveform patterns and alterations in programming parameters and energy delivery), new neural targets, and improved lead placement; neurostimulation has improved in its ability to treat axial LBP.^{1,13–15} Here, we will provide an up-to-date narrative review of the scientific evidence of neurostimulation therapies with a focus on the management of chronic axial LBP.

Traditional Low-frequency Paresthesia-based SCS

Traditional low-frequency (ie frequency ≤ 1000 Hz) paresthesia based SCS has been shown to be clinically effective for many pain conditions including PLPS. However, the ability of SCS to successfully manage chronic axial LBP has been associated with mixed results and limited efficacy; often with declining efficacy over time.^{8,16} A recent systematic review utilizing the Cochrane risk of bias tool and GRADE system demonstrated very low quality evidence for lowfrequency SCS for the treatment of axial LBP with concomitant leg pain.¹⁶ The North American Spine

464

Society (NASS) recently published evidence-based clinical guidelines for multidisciplinary spine care for the treatment of LBP which determined that there was insufficient evidence to make a recommendation for or against the use of SCS for the treatment of LBP.⁶ However, it must be pointed out that new SCS research has been published that was not included in this NASS review. In addition, recent attention has been placed on SCS explant rates with a major driver being loss of clinical efficacy.¹⁷⁻²⁰ Therefore, it is prudent for physicians and manufacturers to understand the current level of evidence for SCS treatment of chronic axial LBP and to continue SCS advancement prior to large-scale clinical utilization for this indication.

When evaluating the ability of traditional lowfrequency SCS to effectively treat lumbar spine conditions, five randomized controlled trials (RCTs; Table 1) are available that include a traditional low-frequency study arm.^{13,14,21-24} These studies all included patients with axial LBP and leg pain. Three of the five studies provide information specifically on back pain reduction with the traditional low-frequency SCS as a secondary outcome.^{13,22,24} The earliest RCT by Kumar et al^{12,24} evaluating SCS LBP reduction as a secondary outcome demonstrated a significant VAS reduction at six months within the SCS group and between the SCS group and conventional medical management. However, by 24 months the between group difference for axial LBP was absent. The two most recent RCTs providing information on the treatment of axial LBP have demonstrated positive sustained results at 12 months, and in one study at 24 months. In the Kapural et al study¹⁴ the traditional SCS arm demonstrated a significant sustained 44% LBP reduction at 24 months. In the Mekhail et al study,²² the traditional SCS arm demonstrated a 54% reduction in LBP at 12 months. In addition to the RCTs, nonrandomized comparative studies and prospective cohort studies have suggested a positive reduction in axial LBP with SCS treatment.¹⁶ PLPS was a common clinical diagtwo nosis and only of the five papers specifically included individuals with nonsurgical refractory LBP. Therefore, there is limited insight on the ability of traditional low-frequency paresthesia based SCS to effectively treat nonsurgical refractory LBP.

Several modifications to traditional low-frequency paresthesia based SCS have been introduced; attempting to improve axial LBP coverage and treatment (Figure 1 and Table 2).^{15,25} The modifications have included attention to stimulation target zones, lead placement, equipment design, programming algorithms, and electrical parameter modifications.^{26–32} Each modification has been suggested to overcome constraints in either clinical, physiological, or anatomical areas. Theorized anatomical constraints have included the close relationship with the dorsal columns to the dorsal root fibers, the increased thickness of the dorsal cerebral spinal fluid at T8-T9, and the size of the L1 and L2 dermatomes with their close proximity to the T12 dermatome.¹⁵ Increased stimulation in the T12 dermatome may result in unwanted chest wall coverage.³³ One of the major challenges with traditional low-frequency SCS has been the ability to capture axial LBP with paresthesia coverage. Paresthesia capture was once considered to be of critical importance because initial work demonstrated that superposition of stimulation paresthesias upon a patient's pain topography to be a predictor of successful pain relief.³⁴ However, current SCS research has demonstrated that continuous paresthesia coverage is not required for pain relief with specific programs and frequency settings,^{21,25,35}

Closed-loop SCS

Conventional SCS modalities use preset programs with regard to frequency, pulse-width, and amplitude which are used to modulate pain processing (Figure 2). These conventional SCS modalities are open-loop systems which require the operator (eg patient, physician, or device representative) to manually change fixed output stimulator settings. To improve open-loop systems, sensing technology has been employed to alternate between preset programs for specific positions.³⁶ Automatic position adaptive stimulation (Figure 2) provides benefits for pain relief, convenience, and a reduction in daily number of amplitude adjustments in comparison to traditional SCS systems (18.2 adjustments per day vs 30.7 adjustments per day).³⁶

Although automatic position adaptive stimulation with open-loop systems have been helpful, limitations still exist since integrated feedback from the nervous system is not incorporated into stimulation patterns. Recently, in an effort to overcome this limitation, a closed-loop SCS system has been introduced which monitors the physiologic state of the nervous system and modifies SCS parameters in real time based on this input.³⁷ The deployment of

a closed-loop system for medical purposes is not new. This technology has been used extensively in the field of cardiology.³⁷ Previous challenges with closed-loop systems that have limited clinical neuromodulation introduction have included technology restrictions (eg sensing and stimulation through the same lead), hardware design, programming intricacies, and increased energy requirements due to computational, sensing and stimulation duties.³⁸

Learning from previous clinically implemented closedloop systems (Figure 2), the therapeutic effects associated with low-frequency traditional paresthesia-based SCS may be increased. Utilizing a closed-loop SCS system that employed evoked compound action potential (ECAP) sensing, Russo et al^{39,40} examined the ability to effectively treat individuals with lower back and/or leg pain. For this study, patients were trialed and implanted with a closedloop SCS system with integrated feedback control using ECAPs to provide a measure of the response of nerve fibers to stimulation and to automatically maintain desired dorsal column fiber recruitment levels within the therapeutic window. The system consisted of two 12-contact percutaneous leads. Preliminary results were published for the three- and six-month outcomes. Fifty-seven percent of patients had PLPS and the primary region of pain was the lower back for approximately 72% of patients. Fortynine patients completed the end of trial assessment and 36 patients elected to move forward to implantation. At six months, 85.7% of patients reported LBP relief >50%. Due to its short-term success, the study was continued further to provide 12-month outcomes which were published in 2019.⁴⁰ The 12-month data demonstrated that 76.9% of patients reported \geq 50% total pain reduction and 56.4% of patients reported ≥80% LBP reduction; therefore, providing clinical evidence to support the use of the closed-loop SCS system not only for neuropathic radicular pain but also for the treatment of axial LBP.⁴⁰

A multicenter double-blind RCT enrolling 134 patients examining the ability of a closed-loop SCS system to treat chronic back and leg pain further provided evidence of safety and efficacy.²² Patients were randomized either to an ECAP controlled closed-loop or fixed output open-loop SCS systems. The baseline demographics demonstrated that 57% to 61% of the patients had a diagnosis of PLPS and 58% to 61% of the patients had previous back surgery depending on the treatment arm. The primary analysis was at three months and follow-up was continued to 12 months (prespecified analysis). At three months, 82.3% of the closed-loop SCS

Table I	Randomized	Controlled	Trials of	Traditional	SCS for	Lumbar Back Pain	
---------	------------	------------	-----------	-------------	---------	------------------	--

Author	Type of Study	Number of Trial Patients n	Successful Trials	Trial Patients Progressing to Implant	Patients with PLPS	Patients without Previous Back Surgery	Range of Follow- Up	Pain Pattern	Pertinent Inclusion and Exclusion Criteria
Kumar et al 2007 ²⁴ and 2008 ¹²	Traditional vs CMM	100	Traditional: 83%	89%	100%	0%	6–24 months	Radicular >axial	Inclusion: PLPS; predominant leg pain. Exclusion: predominant back pain.
Kapural et al 2015 ¹³ and 2016 ¹⁴	Traditional vs 10-kHz HF	198	Overall: 90% 10-kHz HF: 93% Traditional: 88%	86%	77%	13%	12–24 months	Radicular and/or axial	Inclusion: trunk/limb pain refractory to CMM for >3 months.
De Andres 2017 ²³	Traditional vs 10-kHz HF	60	Overall: 92% 10-kHz HF: 90% Traditional: 94%	92%	100%	0%	12 months	Radicular	Inclusion: PLPS; pain refractory to CMM for >6 months Exclusion: mechanical low back pain; coexisting chronic pain or neurological disease.
Deer et al 2018 ²¹	Traditional vs burst	121	Overall: 93% Randomized after trial	83%	42%	Not documented	12 to 24 weeks	Radicular and/or axial	Inclusion: trunk and/or limb pain >60 mm on VAS during 7-day pain diary.
Mekhail et al 2020 ²²	Traditional vs closed- loop	134	Overall: 90% Closed-loop: 94% Traditional: 85%	86%	59%	Not documented	12 months	Radicular and/or axial	Inclusion: trunk/limb pain refractory to CMM; >60 mm on VAS

Note: All percentages rounded to whole number unless <1%.

Abbreviations: SCS, spinal cord stimulation; CMM, conventional medical management; 10-kHz HF, 10 kHz high-frequency therapy; VAS, visual analog scale; PLPS, postlaminectomy pain syndrome (ie FBSS, failed back surgery syndrome); NRS, numeric rating scale; ECAP, evoked compound action potential; LM, lead migration; LOP, loss of paresthesia; PIS, pain at incision site; INF, infection; ULP, unsuccessful lead placement; PGPP, pulse generator pocket pain; MS/C, muscle spasm/cramps; USL, unintended stimulation location.

system patients had achieved the primary outcome of \geq 50% reduction in overall back and leg pain with no increase in medications (Table 1). This trend was continued at 12 months, 83.1% of the patients with the closed-loop system and 61% of the patients with the traditional

SCS system met the primary outcome. When specifically examining LBP scores, 81% of the closed-loop patients and 57% of the open-loop patients achieved \geq 50% reduction in back at three months. When following these patients to 12 months, 80% of the closed-loop patients

Average Percent	Pain Reduction for S	CS at Last Follow-up	Back Pain	Patient	scs	Comments	
Back	Leg	Overall	Responder Rates (≥50% Pain Reduction)	Satisfaction	Complications		
13%	46%	Not documented	Not documented	93%	LM (14%) LOP (12%) PIS (12%) INF (10%)	At 24 months, PLPS patients reported sustained pain relief, improvements with functional capacity and health-related quality of life, and satisfaction with treatment.	
10-kHz HF: 67% Traditional: 44%	10-kHz HF: 70% Traditional: 49%	Not documented	Back pain – 3 months: 10-kHz HF: 84% Traditional: 44% 6 months: 10-kHz HF: 76% Traditional: 52% 12 months: 10-kHz HF: 79% Traditional: 51% 24 months: 10-kHz HF: 77% Traditional: 49%	10-kHz HF: 83% Traditional: 79%	PIS (12%) INF (7%) LM (5%)	Results indicate that 10-kHz high-frequency therapy is superior to traditional SCS.	
Not documented	Not documented	10-kHz HF: 24% Traditional: 19%	Not documented	Not documented	LM (13%)	Results indicate that PLPS can be treated with similar effectiveness by traditional or 10-kHz therapy.	
Burst 5.7 mm less than traditional on 100 mm VAS	Burst 4.7 mm less than traditional on 100 mm VAS	Burst 5.1 mm less than traditional on 100 mm VAS	Not documented	89%	ULP (0.6%) PIS (0.6%)	Purpose of study was to determine safety and efficacy of a device capable of delivering tonic or burst stimulation.	
Closed-loop: 69% Traditional: 54%	Closed-loop: 73% Traditional: 62%	Closed-loop: 72% Traditional: 56%	Back pain – 3 months: Closed-loop: 81% Traditional: 57% 12 months: Closed-loop: 80% Traditional: 58%	Not documented	LM (7%) PGPP (4%) MS/C (2%) USL (2%) INF (1%)	Evidence indicates superiority of closed-loop (ECAP controlled) system.	

and 58% of the open-loop patients maintained \geq 50% back pain reduction. A hypothesized reason for improvement in pain control seen with the closed-loop system was increased time within the therapeutic window for the individuals in the closed-loop group. In conclusion, the use of a closed-loop SCS system allows for optimal therapeutic effects to be maintained without being diminished by the patient's posture or physical activity. Although further research with longer-term outcome data and a large cohort is necessary, a closed-loop system that



Figure I Radiographical images of implanted percutaneous leads. (A) Anterior/posterior view showing the left lead positioned at the top of T8 with the right lead positioned at the middle of T8. The staggered configuration provides overlap of the T8–T9 disc space and T9–10-disc space corresponding to the targeted dermatomes for low back and leg pain. (B) Lateral view indicating the leads are correctly positioned in the posterior epidural space.

is capable of automatically monitoring and adapting stimulation parameters may prove to be a more efficacious treatment option for those suffering from axial symptoms.

High-density or High-dose SCS

Within the last few years, clinicians have been altering parameters that influence electrical charge delivery to find efficacious programming parameters that provide paresthesia-free or -reduced SCS pain relief within frequency ranges ≤1200 Hz. High-density or highdose (HD) SCS is a proposed programming method to improve pain control with SCS through alterations in pulse width and frequency in order to provide a higher charge delivery over a specific period of time (ie higher charge per second).⁴¹ A functional MRI study suggested that HD SCS may modulate the descending pain modulatory system in the brain and brainstem regions in humans with PLPS resulting in supraspinal effects and subsequent pain relief.⁴² To date, no RCTs exist examining the ability of HD to provide effective long-term axial LBP (Table 3). Studies have predominantly consisted of retrospective and prospective case series and often include patients with decreasing efficacy with traditional SCS settings that either have neuropathic pain, complex regional pain syndrome (CRPS) or PLPS.43-47

The earlier studies with HD suggested limited efficacy in treating PLPS patients with LBP and leg pain (Table 3).^{25,43,45} In two retrospective case series with a minimum of 12 months follow-up, HD programming was shown to have positive effects on pain scores.^{25,43} However, when specifically analyzing PLPS patients with low back and leg pain, response rates and pain reduction were not as great as those seen with individuals suffering from other neuropathic pain states (eg CRPS). Sweet et al⁴⁵ examined 15 patients responding to conventional SCS in a prospective case series that switched patients to subthreshold HD. Unfortunately, only four of the 15 patients responded to subthreshold HD programming. However, in this subgroup HD programming resulted in improvement in pain control compared to sham and lower levels of focus on the pain compared to conventional programming. Two small retrospective case series with 12 months follow-up also demonstrated limited efficacy in patients with PLPS.^{25,43} The responder rates based on each study definition for PLPS was 22-33% and in one series many patients aborted HD programming.

Recently, a prospective open label study of 44 SCS naïve patients with PLPS suffering with chronic back and leg pain further examined HD parameters (1000 Hz and 90 μ s or 300 Hz and 800 μ s) for three months and

Table 2 Modifications to Improve Traditional SCS Low Back Coverage

Modification	Study	Pertinent Inclusion and Exclusion Criteria	Findings	Warranted Future Research
Stimulation target zoi	ne			
Target region: T9– T10	Sharan et al 2002 ²⁶	Inclusion: previous lumbar spine surgery.	Cathode positioned at T9–T10; 6–12 months postop, most effective cathode position was at T10 and lower.	Further examination of optimal lead position.
Target region: T10–T11; modified anchoring methods	Mironer et al 2008 ²⁷	Inclusion: axial and/or radicular symptoms maintained after >2 years of SCS treatment.	The use of a single, percutaneous, octad lead with midline anchoring allowed for better coverage of axial and lower extremity pain.	Further examination of optimal lead selection, lead position, and parameters of stimulation.
Target region: T9– T10; non-neuronal cells in the nervous system	Ruiz-Sauri 2019 ⁷²	Dissection of posterior spinal cord segments from 11 human cadavers.	Glial to neuronal ratio in posterior gray matter within the T8– T11 vertebral region ranges from 11:1 to 13:1. At T9– T10, population of oligodendrocytes is significantly larger than any other segment.	Further clinical investigation of the role of glial cells in the development and maintenance of chronic pain. The ability of SCS to modulate glial cells and their relationship to pain control.
Lead design/stimulation	on parameters			
Anode-Cathode Configuration	Holsheimer and Wesselink 1997 ²⁸	Computer modeling	Widest area of coverage found with bipolar or tripolar configuration on single electrode.	Further examination of optimal electrode configuration for low back coverage
Multiple electrode paddle lead	Barolat et al 2001 ²⁹	Inclusion: Axial symptoms ≥ radicular symptoms. Exclusion: PLPS patients excluded if leg pain >back pain.	Multiple electrode paddle lead was found to decrease leg and back pain, while improving quality of life at 6 and 12 months compared to baseline.	Randomized controlled trial with large cohort comparing the efficacy of paddle leads vs percutaneous leads.
Multiple electrodes/ leads	North et al 2005 ³⁰	Inclusion: PLPS with back pain >leg pain.	There was no advantage in pain coverage using two 4-electrode leads vs a single 4-electrode lead.	Further examination of optimal number of leads and lead spacing.
Laminectomy vs percutaneous electrodes	North et al 2006 ³¹	Inclusion: PLPS with back pain >leg pain.	2x8 paddle lead provided no benefit over a single 1x4 percutaneous lead for the treatment of axial pain and resulted in increased power requirements.	Longer-term study comparing the effectiveness of laminectomy vs percutaneous electrode with new programming and SCS system designs.
Sensor-driven position-adaptive system	Schultz et al 2012 ³⁶	Inclusion: axial and/or radicular symptoms; PLPS patients included.	Stimulation parameters are altered as body position changes.	Further examination of sensor-driven position-adaptive systems
Anatomically guided 3D neural targeting	Veizi et al 2017 ³²	Inclusion: PLPS with axia and/or radicular symptoms.	SCS therapy with 3D neural targeting algorithm was more effective at targeting axial symptoms than traditional SCS stimulation therapy.	Further examination of 3D neural targeting for axial low back pain with and without previous lumbar spine surgery.
ECAP controlled closed-loop system	Mekhail et al 2020 ²²	Inclusion: axial and/or radicular symptoms.	At 12 months, 83.1% of patients in the closed-loop group achieved ≥50% decrease in axial/radicular symptoms vs 61.0% in the open-loop group.	Development of methods to reduce the needed energy input to maintain the ECAP controlled system.

Note: Adapted from Provenzano DA, Rebman J, Kuhel C, Trenz H, Kilgore J. The efficacy of high-density spinal cord stimulation among trial, implant, and conversion patients: A retrospective case series. Neuromodulation. 2017;20(7):654–660. Copyright 2017, Wolters Kluwer Health, Inc.²⁵

Abbreviations: PLPS, postlaminectomy pain syndrome; SCS, spinal cord stimulation; ECAP, evoked compound action potential.



Figure 2 A schematic model depicting the difference in conventional SCS systems compared to closed-loop systems. The conventional system has set stimulation parameters. The closed-loop system alters stimulation parameters depending on the activity, body position, and physiological state of the patient in real time using sensor feedback.

provided insight on the ability to treat LBP.⁴⁸ HD programming was found to have a significant reduction in overall pain (NRS; 7.5 to 3.8; P<0.01), LBP (7.2 to 3.4; P<0.01), leg pain (7.2 to 3.1; P<0.01) with improvements in functional status (ie Oswestry Disability Index).

Recently, the durability of HD programming was further evaluated in a 12-month prospective single arm

multicenter study evaluating individuals with chronic intractable low back and leg pain using a standardized approach to HD programming with 91 patients (103 patients were initially implanted) available at that 12-month follow-up.⁴⁷ At the 12-month visit, there was a statistically significant reduction in overall pain (56.7%), LBP (51.9%) and leg pain (63.7%) compared to baseline.

In summary, the research to date on HD programming has demonstrated mixed results with regard to the treatment of axial LBP; especially in individuals with PLPS. In addition, its ability to improve pain control in those that have lost efficacy with traditional SCS has been questioned.⁴⁴ However, the recent single arm prospective study with 12-month data has suggested the sustained ability to treat axial LBP.47 Long-term research with a control arm is needed to confirm the results of the longer-term prospective trials. Furthermore, it has been hypothesized that responder rates are improved when mean pulse densities and charge per second are increased.43 However, it must be remembered that alterations in frequency and pulse-widths within traditional SCS settings alone without consideration of charge per second or pulse density have been shown to positively influence sensory thresholds (ie mechanical detection and pain pressure thresholds), therapeutic range, paresthesia coverage, and patient sensation/satisfaction scores.^{49,50} Dosing studies have also demonstrated that lower frequencies utilizing lower charge per second may provide equivalent pain relief when pulse-width and amplitude settings are optimized.⁵¹ Specifically, the PROCO RCT investigated the effects of rate on analgesia in kilohertz frequency (1-10 kHz) in individuals with low back pain with or without equal or lesser leg pain. When pulse width and amplitude were adjusted to optimize subperception SCS therapy while stimulating at the previously identified stimulation location, all tested frequencies provided equivalent pain relief. Therefore, titrating to a specific charge per unit time may not be needed or indicated. No research to date exists regarding the ability of HD to treat nonsurgical refractory LBP.

Burst Spinal Cord Stimulation

This section will discuss burst stimulation defined by groups of five spikes of energy delivered at 500 Hz with a pulse width of one millisecond and one millisecond interspike intervals at subthreshold amplitudes followed by a passive discharge phase delivered over five milliseconds.⁵² Burst stimulation was first implemented for the treatment of chronic neuropathic pain in 2010.⁵³ Burst SCS may induce distinct patterns of modulation of neurons in the dorsal horn and dorsal column nuclei and brain activation.⁵⁴ To date, limited research is available to evaluate the efficacy of burst SCS in the treatment of axial and nonsurgical refractory LBP.^{16,55,56}

Currently, the majority of the research published on burst SCS involves heterogeneous subject populations which makes it challenging to evaluate its efficacy for the treatment of predominant axial LBP.^{8,16,55} To date, only a nonrandomized comparative study exists evaluating the efficacy of burst SCS in the treatment of predominant axial LBP.⁵⁷ In this comparative study between burst and 10-kHz high-frequency SCS (10-kHz HF), 16 subjects with PLPS were enrolled with predominant LBP (\geq 70% of global pain) and results were presented with a mean post-implant follow-up range of 10–20 months. Both therapies demonstrated a significant decrease from baseline to long-term follow-up in mean LBP intensity with no difference between groups (87.5±17.7% and 54.9±44% reduction for burst SCS and 10-kHz HF, respectively).⁵⁷

In the largest RCT on burst SCS (SUNBURST TRIAL) the effectiveness of burst stimulation in comparison to tonic stimulation was evaluated in 100 subjects (all candidates for a currently approved neurostimulation device) that were randomized to either treatment and followed for up to one year.²¹ In this study there was a heterogeneous subject population that included patients with back pain, leg pain, nonsurgical refractory low back pain (ie no history of previous back surgery) and other non-spine conditions (eg CRPS). PLPS was diagnosed in 41.8% of patients. Burst SCS was preferred by 70.8% of patients. Responders to treatment were classified as individuals achieving $\geq 30\%$ reduction in overall daily VAS score from baseline. Based on the intention-to-treat analysis, the estimated difference in the overall VAS score between burst and tonic stimulation was -5.1 mm on a 0-100 mm VAS scale. When evaluating the number of patients that had clinically significant reductions in pain $(\geq 30\%)$, 60% of the burst SCS patient and 51% of the tonic patients met or exceeded this cutoff (Table 1). Unfortunately, the study did not allow for detailed evaluation of the ability of burst SCS to treat predominantly axial LBP based on the study population and data collected.

Recently, a systematic review and meta-analysis of RCTs and prospective cohort studies was conducted comparing SCS waveforms and programming for treating chronic LBP predominantly in individuals with PLPS.⁵⁶ Specifically, six studies were evaluated comparing burst SCS vs traditional SCS and two studies comparing burst SCS vs 10-kHz HF SCS. Although many studies included a high risk of bias, a meta-analysis of five studies comparing for traditional SCS vs traditional SCS revealed pooled super-iority of burst over traditional for chronic LBP reduction.⁵⁶

nts Pare Preid Surr Bac 0% Surr 0% Not Not	ients hout k gery rred	Range of Follow- up B weeks (four two- week periods) 12 nonths 12	Neuropathic Pain Pattern Axial/radicular Radicular Axial/radicular Axial/radicular	Pertinent Inclusion & Exclusion & Criteria Criteria Criteria Inclusion: PLPS treated initially with traditional SCS. Exclusion: patients week HD trial. week HD trial. hnclusion: patients with unsuccessful traditional SCS treatment. Exclusion: Patients with <3 months of traditional SCS treatment. Exclusion: minimum of one year HD	Average I Reduction Not reported Not Not reported	Eercent Pai Leg Not reported 51% Not reported	n Overall Not reported Trial	Patient Satisfaction Not Not Not Not Not	Summary of Results Upon Completion of the trial, all patients requested the option to continue with subthreshold HD Subthresho
			extremity pain	of one year TU treatment follow- up.	reported	reported	patients 54% HD: 72% Conversion patients 50%	reported	surrering roun CRPS responded more frequently to HD programming than individuals
							HD: 43%		with PLP

472

submit your manuscript | www.dovepress.com
DovePress

Dovepress

% 0% 12 Axial and/or Inclusion: PLPS with 34% 39% Not Not Conversion months radicular unsuccessful nusccessful reported reported from radicular traditional SCS traditional SCS reported reported from restment. Exclusion: life An effective respectancy <l td="" year<=""> An effective an effective and leg pain. respectancy <l td=""> Pain relief is relieve back Pain relief is relieve back Pain relief is relieve back Pain relief is rest L2 Pain relief is relieve back</l></l>	0% 3 Axial and Inclusion: PLPS 53% 54% 84% Patients that used 1000 Hz months Radicular Exclusion: prior exposure to 90 µ pulse- 90 µ pulse- inclusion exposure to exposure to exposure to 90 µ pulse- width inclusion exposure to exposure to exposure to exclusion therapy. exclusion therapy. inclusion exposure to exposure to exposure to exclusion therapy. exclusion therapy. inclusion exposure to exposure to exposure to exclusion therapy. exclusion therapy. inclusion exposure to exposure to exposure to exclusion therapy. exclusion therapy. inclusion exposure to exposure to exposure to exclusion therapy. inclusion exposure to exposure to exclusion therapy. exclusion therapy. inclusion exposure to exposure to exclusion therapy. exclusion therapy.	Not12Axial/radicularInclusion: baseline52%64%57%85%Study resultsreportedmonthspain of ≥ 50mmPain
l/or Inclus unsuc treatri Excluu expec	Exclus expose stimul	icular Inclus pain c VAS; V VAS; v able t medic Exclus neuro therar
Axial and radicular	Axial and Radicular	Axial/radi
12 months	3 months	12 months
%0	%0	Not reported
8001	%001	45%
28	84	-
Prospective Conversion from traditional to HD SCS	Prospective Conversion from traditional to HD SCS	Prospective HD SCS
De Jaeger et al 2020 ⁴⁶	Benyamin et al 2020 ⁴⁸	Hatheway et al 2020 ⁴⁷

Dovepress

473

Further examination of burst SCS for the treatment of axial LBP and nonsurgical refractory LBP with longitudinal follow-up is warranted. Studies would benefit from narrowing patient selection to provide optimal insight into the treatment of axial LBP. Recently, additional efforts have been made to improve our understanding of the deployment of burst SCS in the treatment of LBP including the evaluation of lead placement (anatomical placement vs paresthesia mapping), optimization of dosing, and stimulation parameters.^{58–61}

10-kHz High-frequency Therapy

Ten kilohertz high-frequency therapy utilizes a higher frequency (10-kHz) than that of traditional stimulation as well as substantially shortened pulse-width (30 µs) to elicit pain relief without the sensation of paresthesia.⁶² In addition, the pain relief associated with 10-kHz HF therapy is paresthesia independent and therefore leads are placed based on anatomical landmarks.³⁵ Previous research has suggested that frequency, as well as other parameters including pulse width and amplitude, is integral in pain reduction.^{50,51} A randomized crossover study evaluating subthreshold frequencies of 1200 Hz, 3030 Hz, and 5882 Hz, demonstrated that a frequency of 5882 Hz produced significant axial LBP relief in comparison to lower frequencies and sham SCS.⁶³ As with other modes of SCS, the underlying mode of action is still being elicited. Basic science research has suggested that 10-kHz stimulation suppresses spontaneous activity in the dorsal horn and the preferential blockade of large diameter fibers with concomitant recruitment of medium and small fibers.⁶⁴⁻⁶⁶ Furthermore, structural volumetric changes in areas of the brain involved in chronic pain processing have been shown to be influenced.⁶⁷ Specifically, significant decreases were found in the volume in the left and right hippocampus over time.

A multicenter RCT with 24-month follow-up and recent systematic review utilizing the Cochrane risk of biased TOOL and GRADE system provide evidence that 10-kHz SCS, when compared to low-frequency SCS, is more effective in patients with axial LBP and concurrent leg pain.^{13,14,16} In the 24-month multicenter RCT conducted, 76.5% of patients were responders to 10-kHz HF therapy for LBP compared to 49.3% for traditional SCS.¹⁴ In addition, 60% of patients treated with 10-kHz HF therapy were satisfied with their reduction in pain at 24 months vs only 40.4% of traditional SCS patients.¹⁴ In this study 87% of patient had previous back surgery with baseline back scores >7 on the

VAS. The recent systematic review demonstrated lowquality evidence that 10-kHz is effective beyond six months for axial LBP reduction in patients with predominantly axial spine pain and those with mixed axial low back and leg pain.¹⁶ A non-industry funded SCS study examining 10kHz HF to traditional SCS demonstrated no difference in global pain reduction at one year.⁶⁸ However, back pain scores were not reported separately.⁶⁸

Initial research has investigated the ability of 10-kHz HF therapy to treat nonsurgical refractory LBP.⁶⁹ In a prospective open label study of individuals with chronic axial LBP without previous spinal surgery, 21 patients were examined. 95% of the patients went on to implantation and 17 patients were followed for 36 months. Significant reduction in pain (79 ± 12 mm to 10 ± 12 mm), improvement in functional status, and reduction in opioid utilization were demonstrated. A subanalysis of pooled data from two prospective studies evaluating 10-kHz HF therapy for the treatment of nonsurgical refractory back pain demonstrated a 70% reduction in average back pain and improvement in Oswestry disability index (ODI) with reduced opioid consumption at 12 months in the combined cohort.¹

Differential Target Multiplexed SCS

Differential target multiplexed SCS is a novel programming approach based on evidence from preclinical studies.^{64,70} Differential target multiplexed stimulation uses multiple electrical signals that differ from one another in electrical wave parameters which has been suggested to stimulate multiple target tissues including pain fibers in the dorsal column and modulate glial cells in the spinal cord.⁷⁰ Differential target multiplexed stimulation aims to modulate chronic pain-related biological processes back toward their non-pain state.⁷¹ Basic science research has indicated that glial cells outnumber neurons in the spinal cord 20:1, are electrically excitable, and have the ability to modulate ascending pain pathways.^{72,73} This evidence allowed researchers to hypothesize that SCS parameters can be adjusted to differentially target neurons and glial cells in order to maximize the efficacy of treatment.⁶⁴

A prospective multicenter feasibility dual trial study was conducted with 25 patients with chronic intractable back pain with or without leg pain that underwent a threeto-five-day SCS trial using traditional stimulation followed by a three-to-five-day trial using differential target multiplexed stimulation. The trial periods were separated by a 24-h washout period.⁷⁰ Differential target multiplexed therapy resulted in a \geq 50% back pain relief in 80% of the patients while traditional SCS provided \geq 50% relief in 50% of the patients. Specifically, differential target multiplexed therapy resulted in a 67% reduction in LBP compared to baseline while traditional SCS resulted in a 43% reduction from baseline (*P*<0.0001).

Further published studies evaluating the efficacy of differential target multiplexed programming in a RCT setting with longer-term follow-up (minimum of 12 to 24 months) and refinements of the study population are needed to evaluate the long-term effectiveness in treating axial LBP and nonsurgical LBP.

Dorsal Root Ganglion Stimulation

Preliminary research has examined the utilization of DRG stimulation to treat chronic low back pain.^{74–78} The DRG contains the primary sensory neurons that transmit afferent nociception.⁷⁴ Currently, DRG stimulation is FDA approved for the treatment of complex regional pain syndrome of the lower limbs. To date, four case series with a total of 60 patients have examined the effectiveness of DRG specifically for the treatment of low back pain with 12-month follow-up available in some studies.^{75–78} Dorsal root ganglions that were specifically stimulated included T12,⁷⁵ and L1–L5^{76–78} In these case series, DRG stimulation has been shown to improve pain levels, physical and mental functioning, disability, and quality of life. Future well-developed RCTs with long-term follow-up are needed to confirm these initial findings including safety outcomes.^{79,80} Furthermore, the optimal DRG stimulation level for the treatment of chronic axial lumbar pain needs to be further refined.

Peripheral Nerve Stimulation (Medial Branch Nerve Innervation to the Multifidus Muscle)

Forms of electrical stimulation for the treatment of axial LBP include other modalities besides those that target the dorsal columns. Stimulation of the medial branch nerves innervating the multifidus muscle is one type of non-spinal cord stimulation that has shown promising results.⁸¹

The multifidus muscle extends lengthwise on either side of the spinal column from the cervical region to the sacral region and acts as an extensor and back stabilizer. Lumbar multifidus muscles generate large forces over short distances for intersegmental control.^{81,82} Functional instability due to weakness of the lumbar multifidus muscle is a significant contributor to axial LBP.⁸² Physiotherapy is often used as a noninvasive treatment option to strengthen the multifidus muscle and reduce axial symptoms; however, long-term relief may be challenging secondary to compliance. Multifidus stimulation is conducted using percutaneous electrodes implanted over the relevant medial branches. Efficacy of the treatment is associated with thickening of the multifidus muscle.⁸³ A study conducted by Sions et al⁸³ indicated an average of 20.38% thickening of the lumbar multifidus muscles utilizing multifidus stimulation therapy. Deckers et al⁸¹ conducted a study using multifidus stimulation on 53 subjects with chronic mechanical LBP. Leads were placed bilaterally over the L2 medial branch nerves at the junction of the transverse process and superior articular process. After 90 days, 58% of subjects had \geq 2-point reduction in NRS pain scores. At 12 months, 57% had ≥2-point reduction in NRS pain scores and 60% had \geq 10-point improvement in ODI. Eighty-one percent of patients in this study were satisfied with their treatment after one year. Cohen et al⁸⁴ performed a case series of nine subjects using temporary (one-month) percutaneous multifidus stimulation for chronic LBP. Using ultrasound guidance, electrodes were placed 1-2 cm posterior to the lamina at the center of the patients reported pain to target the medial branch of the dorsal primary ramus. At one month, 67% of patients reported \geq 50% reduction in LBP. Among responders, an 80% reduction in LBP was achieved. Pain reduction was maintained at 80% of baseline pain for three months after lead removal and gradually increased to 60% of baseline pain at six months after lead removal.

Stimulation methods that target painful areas more directly than conventional SCS modalities show promising results. Although currently there is not overwhelming evidence to support the use of these treatment modalities. Multifidus stimulation may be effective at treating symptoms related to muscle weakness in the lumbar region; however, symptoms relating to neuropathic pain will presumably remain. Additional studies looking at the use of multifidus stimulation for chronic LBP are needed.

Conclusion

Spinal cord stimulation and peripheral nerve stimulation have been studied extensively for the last half century. Recently, attention has been placed on the ability of neurostimulation to treat axial LBP and refractory nonsurgical LBP. Current evidence of efficacy specifically restricted to these conditions is limited. A majority of

475

SCS research evaluating treatment efficacy for axial LBP include industry sponsored studies with heterogeneous patient populations often with radicular symptoms and history of prior back surgery. Traditional SCS has demonstrated long-term efficacy for treating radicular symptoms, but limited efficacy for the treatment of axial LBP. Multiple enhancements and modifications in traditional SCS including our understanding of stimulation target zones, electrode placement, electrical parameters, and equipment design modifications may result in improved outcomes. Furthermore, advancements in novel programming, electrical signaling, energy delivery, waveforms and technologies have allowed for the ability to treat pain in a paresthesia independent manner. Neurostimulation therapies that do not target the spinal cord may also be associated with significant axial LBP reduction.

Finally, the minimal invasiveness of neurostimulation remains a compelling reason for patients to seek this treatment option for the treatment of axial low back pain. Invasive surgical methods (eg fusion) that alter the anatomy of the spine with considerable rates of failure and high adverse event rates are often considered before neurostimulation. If neurostimulation is shown to demonstrate long-term effectiveness in appropriately designed RCTs with low complication and explant rates, then neurostimulation therapies may move up in the treatment algorithm for chronic axial LBP and refractory nonsurgical LBP.

Abbreviations

SCS, spinal cord stimulation; NASS, North American Spine Society; LBP, low back pain; HD, high density; 10-kHz HF, 10 kHz high-frequency therapy; CMM, conservative medical management; QALY, quality adjusted life year; IPG, internal pulse generator; FBSS, failed back surgery syndrome; PLPS, post laminectomy pain syndrome.

Funding

No funding was obtained for the writing of this manuscript.

Disclosure

Dr Provenzano has consulted for Abbott, Avanos, Boston Scientific, Esteve, Medtronic, Nevro, Stimgenics and Heron; has received research support from Avanos, Medtronic, Nevro, Stimgenics, and Abbott; and is an owner of Western PA Surgical Center. Dr Hanes has consulted for Medtronic, Abbott and Boston Scientific; has received research support from Boston Scientific; and is an owner of Jax Spine & Pain Centers. The authors report no other potential conflicts of interest in this work.

References

- Al-Kaisy A, Van Buyten JP, Kapural L, et al. 10 kHz spinal cord stimulation for the treatment of non-surgical refractory back pain: subanalysis of pooled data from two prospective studies. *Anaesthesia*. 2020;75(6):775–784. doi:10.1111/anae.15036
- Sinclair C, Verrills P, Barnard A. A review of spinal cord stimulation systems for chronic pain. J Pain Res. 2016;9:481–492. doi:10.2147/ JPR.S108884
- Lamer TJ. Global, regional, and national incidence, prevalence, and years lived with disability for 354 diseases and injuries for 195 countries and territories, 1990-2017: a systematic analysis for the Global Burden of Disease Study 2017. *Lancet.* 2018;392(1015 9):1789–1858.
- Dagenais S, Caro J, Haldeman S. A systematic review of low back pain cost of illness studies in the United States and internationally. *Spine J.* 2008;8(1):8–20. doi:10.1016/j.spinee.2007.10.005
- Andersson GB. Epidemiological features of chronic low-back pain. Lancet. 1999;354(9178):581–585. doi:10.1016/S0140-6736(99)013 12-4
- Kreiner DS, Matz P, Bono CM, et al. Guideline summary review: an evidence-based clinical guideline for the diagnosis and treatment of low back pain. *Spine J.* 2020;20(7):998–1024. doi:10.1016/j.spinee.20 20.04.006
- Shealy CN, Taslitz N, Mortimer JT, Becker DP. Electrical inhibition of pain: experimental evaluation. *Anesth Analg.* 1967;46(3):299–305. doi:10.1213/0000539-196705000-00009
- Deer TR, Grider JS, Lamer TJ, et al. A Systematic Literature Review of Spine Neurostimulation Therapies for the Treatment of Pain. *Pain Med.* 2020;21(7):1421–1432. doi:10.1093/pm/pnz353
- Taylor RS, Ryan J, O'Donnell R, Eldabe S, Kumar K, North RB. The cost-effectiveness of spinal cord stimulation in the treatment of failed back surgery syndrome. *Clin J Pain*. 2010;26(6):463–469. doi:10.1097/AJP.0b013e3181daccec
- Kumar K, Caraway DL, Rizvi S, Bishop S. Current challenges in spinal cord stimulation. *Neuromodulation*. 2014;17(Suppl 1):22–35. doi:10.1111/ner.12172
- Palmer N, Guan Z, Chai NC. Spinal Cord Stimulation for Failed Back Surgery Syndrome – Patient Selection Considerations. *Translat Perioperative Pain Med.* 2019;6(3):81–90.
- Kumar K, Taylor RS, Jacques L, et al. The effects of spinal cord stimulation in neuropathic pain are sustained: a 24-month follow-up of the prospective randomized controlled multicenter trial of the effectiveness of spinal cord stimulation. *Neurosurgery*. 2008;63 (4):762–770. doi:10.1227/01.NEU.0000325731.46702.D9
- Kapural L, Yu C, Doust MW, et al. Novel 10-kHz High-frequency therapy (HF10 therapy) Is superior to traditional low-frequency spinal cord stimulation for the treatment of chronic back and leg pain: the SENZA-RCT randomized controlled trial. *Anesthesiology*. 2015;123(4):851–860. doi:10.1097/ALN.000000000000774
- 14. Kapural L, Yu C, Doust MW, et al. Comparison of 10-kHz high-frequency and traditional low-frequency spinal cord stimulation for the treatment of chronic back and leg pain: 24-month results from a multicenter, randomized, controlled pivotal trial. *Neurosurgery*. 2016;79(5):667–677. doi:10.1227/NEU.000000000001418

- Provenzano DA, Amirdelfan K, Kapural L, Sitzman BT. Evidence gaps in the use of spinal cord stimulation for treating chronic spine conditions. *Spine*. 2017;42:S80–S92. doi:10.1097/BRS.000000000002184
- 16. Conger A, Sperry BP, Cheney CW, et al. The effectiveness of spinal cord stimulation for the treatment of axial low back pain: A systematic review with narrative synthesis. *Pain Med.* 2020;21 (11):2699–2712. doi:10.1093/pm/pnaa142
- Al-Kaisy A, Royds J, Al-Kaisy O, et al. Explant rates of electrical neuromodulation devices in 1177 patients in a single center over an 11-year period. *Reg Anesth Pain Med.* 2020;45(11):883–890. doi:10.1136/rapm-2020-101681
- Van Buyten JP, Wille F, Smet I, et al. Therapy-related explants after spinal cord stimulation: results of an international retrospective chart review study. *Neuromodulation*. 2017;20(7):642–649. doi:10.1111/ner.12642
- Hayek SM, Veizi E, Hanes M. Treatmentlimiting complications of percutaneous spinal cord stimulator implants: A review of eight years of experience from an academic center database. *Neuromodulation*. 2015;18(7):603–609. doi:10.1111/ner.12312
- Pope JE, Deer TR, Falowski S, et al. Multicenter retrospective study of neurostimulation with exit of therapy by explant. *Neuromodulation*. 2017;20(6):543–552. doi:10.1111/ner.12634
- Deer T, Slavin KV, Amirdelfan K, et al. Success using neuromodulation with BURST (SUNBURST) study: results from a prospective, randomized controlled trial using a novel burst waveform. *Neuromodulation*. 2018;21(1):56–66. doi:10.1111/ner.12698
- Mekhail N, Levy RM, Deer TR, et al. Long-term safety and efficacy of closed-loop spinal cord stimulation to treat chronic back and leg pain (Evoke): a double-blind, randomised, controlled trial. *Lancet Neurology*. 2020;19(2):123–134. doi:10.1016/S1474-4422(19)30414-4
- 23. De Andres J, Monsalve-Dolz V, Fabregat-Cid G, et al. Prospective, randomized blind effect-on-outcome study of conventional vs high-frequency spinal cord stimulation in patients with pain and disability due to failed back surgery syndrome. *Pain Med.* 2017;18(12):2401–2421.
- 24. Kumar K, Taylor RS, Jacques L, et al. Spinal cord stimulation versus conventional medical management for neuropathic pain: a multicentre randomised controlled trial in patients with failed back surgery syndrome. *Pain*. 2007;132(1):179–188. doi:10.1016/j.pain.2007.07.028
- 25. Provenzano DA, Rebman J, Kuhel C, Trenz H, Kilgore J. The efficacy of high-density spinal cord stimulation among trial, implant, and conversion patients: A retrospective case series. *Neuromodulation*. 2017;20(7):654–660. doi:10.1111/ner.12612
- 26. Sharan A, Cameron T, Barolat G. Evolving patterns of spinal cord stimulation in patients implanted for intractable low back and leg pain. *Neuromodulation*. 2002;5(3):167–179. doi:10.1046/j.1525-14 03.2002.02027.x
- Mironer YE, Satterthwaite JR, Lewis EM, et al. Efficacy of a single, percutaneous, across midline, Octrode[®] lead using a "midline anchoring" technique in the treatment of chronic low back and/or lower extremity pain: A retrospective study. *Neuromodulation*. 2008;11 (4):286–295. doi:10.1111/j.1525-1403.2008.00178.x
- Holsheimer J, Wesselink WA. Effect of anode-cathode configuration on paresthesia coverage in spinal cord stimulation. *Neurosurgery*. 1997;41(3):654–659. doi:10.1097/00006123-199709000-00030
- Barolat G, Oakley JC, Law JD, North RB, Ketcik B, Sharan A. Epidural spinal cord stimulation with a multiple electrode paddle lead is effective in treating intractable low back pain. *Neuromodulation*. 2001;4(2):59–66. doi:10.1046/j.1525-1403.2001.00059.x
- 30. North RB, Kidd DH, Olin J, et al. Spinal cord stimulation for axial low back pain: a prospective, controlled trial comparing dual with single percutaneous electrodes. *Spine*. 2005;30(12):1412–1418. doi:10.1097/01.brs.0000166502.05449.a8
- 31. North RB, Kidd DH, Olin J, Sieracki JN, Petrucci L. Spinal cord stimulation for axial low back pain: a prospective controlled trial comparing 16-contact insulated electrodes with 4-contact percutaneous electrodes. *Neuromodulation*. 2006;9(1):56–67. doi:10.1111/ j.1525-1403.2006.00043.x

- 32. Veizi E, Hayek SM, North J, et al. Spinal cord stimulation (SCS) with anatomically guided (3D) neural targeting shows superior chronic axial low back pain relief compared to traditional SCS-LUMINA study. *Pain Med.* 2017;18(8):1534–1548. doi:10. 1093/pm/pnw286
- 33. Deer T, Pope J, Hayek S, et al. Neurostimulation for the treatment of axial back pain: a review of mechanisms, techniques, outcomes, and future advances. *Neuromodulation*. 2014;17(Suppl 2):52–68. doi:10. 1111/j.1525-1403.2012.00530.x
- North RB, Ewend MG, Lawton MT, Piantadosi S. Spinal cord stimulation for chronic, intractable pain: superiority of "multi-channel" devices. *Pain*. 1991;44(2):119–130. doi:10.1016/0304-3959(91) 90125-H
- 35. De Carolis G, Paroli M, Tollapi L, et al. Paresthesia-independence: an assessment of technical factors related to 10 kHz paresthesia-free spinal cord stimulation. *Pain Phys.* 2017;20(4):331–341.
- 36. Schultz DM, Webster L, Kosek P, Dar U, Tan Y, Sun M. Sensordriven position-adaptive spinal cord stimulation for chronic pain. *Pain Phys.* 2012;15(1):1–12.
- Denison T, Litt B. Advancing neuromodulation through control systems: a general framework and case study in posture-responsive stimulation. *Neuromodulation*. 2014;17(Suppl 1):48–57. doi:10.1111/ner. 12170
- Sun FT, Morrell MJ. Closed-loop neurostimulation: the clinical experience. *Neurotherapeutics*. 2014;11(3):553–563. doi:10.1007/ s13311-014-0280-3
- 39. Russo M, Cousins MJ, Brooker C, et al. Effective relief of pain and associated symptoms with closed-loop spinal cord stimulation system: preliminary results of the Avalon Study. *Neuromodulation*. 2018;21(1):38–47. doi:10.1111/ner.12684
- Russo M, Brooker C, Cousins MJ, et al. Sustained long-term outcomes with closed-loop spinal cord stimulation: 12-month results of the prospective, multicenter, open-label Avalon Study. *Neurosurgery*. 2020;87(4):E485–e495. doi:10.1093/neuros/nyaa003
- 41. Miller JP, Eldabe S, Buchser E, Johanek LM, Guan Y, Linderoth B. Parameters of spinal cord stimulation and their role in electrical charge delivery: a review. *Neuromodulation*. 2016;19(4):373–384. doi:10.1111/ner.12438
- 42. De Groote S, Goudman L, Peeters R, et al. The influence of high dose spinal cord stimulation on the descending pain modulatory system in patients with failed back surgery syndrome. *NeuroImage: Clinical*. 2019;24:102087. doi:10.1016/j.nicl.2019.102087
- 43. Wille F, Breel JS, Bakker EWP, Hollmann MW. Altering conventional to high density spinal cord stimulation: an energy dose-response relationship in neuropathic pain therapy. *Neuromodulation*. 2017;20(1):71–80. doi:10.1111/ner.12529
- 44. Kapural L, Harandi S. Long-term efficacy of 1–1.2 kHz subthreshold spinal cord stimulation following failed traditional spinal cord stimulation: a retrospective case series. *Reg Anesth Pain Med.* 2019;44 (1):107–110. doi:10.1136/rapm-2018-000003
- 45. Sweet J, Badjatiya A, Tan D, Miller J. Paresthesia-free high-density spinal cord stimulation for postlaminectomy syndrome in a prescreened population: a prospective case series. *Neuromodulation*. 2016;19(3):260–267. doi:10.1111/ner.12357
- 46. De Jaeger M, Goudman L, Brouns R, et al. The long-term response to high-dose spinal cord stimulation in patients with failed back surgery syndrome after conversion from standard spinal cord stimulation: an effectiveness and prediction study. *Neuromodulation*. 2020. doi:10.1111/ner.13138
- 47. Hatheway J, Mangal V, Fishman M, et al. Long-term efficacy of a novel spinal cord stimulation clinical workflow using kilohertz stimulation: twelve-month results from the vectors study. *Neuromodulation*. 2020. doi:10.1111/ner.13324
- Benyamin R, Galan V, Hatheway J, et al. Options: a prospective, open-label study of high-dose spinal cord stimulation in patients with chronic back and leg pain. *Pain Phys.* 2020;23(1):87–98.

- 49. Abejon D, Rueda P, Del Saz J, Arango S, Monzon E, Gilsanz F. Is the introduction of another variable to the strength-duration curve necessary in neurostimulation? *Neuromodulation*. 2015;18(3):182–190. doi:10.1111/ner.12223
- Youn Y, Smith H, Morris B, Argoff C, Pilitsis JG. The Effect of High-Frequency Stimulation on Sensory Thresholds in Chronic Pain Patients. *Stereotact Funct Neurosurg.* 2015;93(5):355–359. doi:10.11 59/000438998
- Thomson SJ, Tavakkolizadeh M, Love-Jones S, et al. Effects of rate on analgesia in kilohertz frequency spinal cord stimulation: results of the PROCO randomized controlled trial. *Neuromodulation*. 2018;21 (1):67–76. doi:10.1111/ner.12746
- De Ridder D, Plazier M, Kamerling N, Menovsky T, Vanneste S. Burst spinal cord stimulation for limb and back pain. *World Neurosurg*. 2013;80(5):642–649. doi:10.1016/j.wneu.2013.01.040
- 53. De Ridder D, Vanneste S, Plazier M, van der Loo E, Menovsky T. Burst spinal cord stimulation: toward paresthesia-free pain suppression. *Neurosurgery*. 2010;66(5):986–990. doi:10.1227/01. NEU.0000368153.44883.B3
- 54. Chakravarthy K, Fishman MA, Zuidema X, Hunter CW, Levy R. Mechanism of action in burst spinal cord stimulation: review and recent advances. *Pain Med.* 2019;20(Suppl Supplement_1):S13–s22. doi:10.1093/pm/pnz073
- 55. Lamer TJ, Moeschler SM, Gazelka HM, Hooten WM, Bendel MA, Murad MH. Spinal stimulation for the treatment of intractable spine and limb pain: a systematic review of RCTs and meta-analysis. *Mayo Clin Proc.* 2019;94(8):1475–1487. doi:10.1016/j.mayocp.2018.12.037
- Karri J, Orhurhu V, Wahezi S, Tang T, Deer T, Abd-Elsayed A. Comparison of spinal cord stimulation waveforms for treating chronic low back pain: systematic review and meta-analysis.. *Pain Phys.* 2020;23 (5):451–460.
- 57. Muhammad S, Roeske S, Chaudhry SR, Kinfe TM. Burst or high-frequency (10 kHz) spinal cord stimulation in failed back surgery syndrome patients with predominant back pain: one year comparative data. *Neuromodulation*. 2017;20(7):661–667. doi:10.1111/ner.12611
- Al-Kaisy A, Baranidharan G, Palmisani S, et al. Comparison of paresthesia mapping to anatomical placement in burst spinal cord stimulation: initial trial results of the prospective, multicenter, randomized, double-blinded, crossover, crisp study. *Neuromodulation*. 2020;23(5):613–619. doi:10.1111/ner.13104
- 59. Schu S, Slotty PJ, Bara G, von Knop M, Edgar D, Vesper J. A prospective, randomised, double-blind, placebo-controlled study to examine the effectiveness of burst spinal cord stimulation patterns for the treatment of failed back surgery syndrome. *Neuromodulation*. 2014;17(5):443–450. doi:10.1111/ner.12197
- 60. Vesper J, Slotty P, Schu S, et al. Burst SCS Microdosing is as efficacious as standard burst scs in treating chronic back and leg pain: results from a randomized controlled trial. *Neuromodulation*. 2019;22(2):190–193. doi:10.1111/ner.12883
- 61. Van Havenbergh T, Vancamp T, Van Looy P, Vanneste S, De Ridder D. Spinal cord stimulation for the treatment of chronic back pain patients: 500-Hz vs. 1000-Hz burst stimulation. *Neuromodulation*. 2015;18(1):9–12. doi:10.1111/ner.12252
- 62. Russo M, Van Buyten J-P. 10-kHz high-frequency SCS therapy: a clinical summary. *Pain Med.* 2015;16(5):934–942. doi:10.1111/pme.12617
- 63. Al-Kaisy A, Palmisani S, Pang D, et al. Prospective, randomized, sham-control, double blind, crossover trial of subthreshold spinal cord stimulation at various kilohertz frequencies in subjects suffering from failed back surgery syndrome (SCS Frequency Study). *Neuromodulation*. 2018;21(5):457–465. doi:10.1111/ ner.12771
- 64. Vallejo R, Kelley CA, Gupta A, Smith WJ, Vallejo A, Cedeño DL. Modulation of neuroglial interactions using differential target multiplexed spinal cord stimulation in an animal model of neuropathic pain. *Mol Pain.* 2020;16:1744806920918057. doi:10.1177/17448 06920918057

- Vallejo R, Bradley K, Kapural L. Spinal cord stimulation in chronic pain: mode of action. *Spine*. 2017;42:S53–S60. doi:10.1097/ BRS.00000000002179
- Arle JE, Mei L, Carlson KW, Shils JL. High-frequency stimulation of dorsal column axons: potential underlying mechanism of paresthesia-free neuropathic pain relief. *Neuromodulation*. 2016;19 (4):385–397. doi:10.1111/ner.12436
- 67. De Groote S, Goudman L, Van Schuerbeek P, et al. Effects of spinal cord stimulation on voxel-based brain morphometry in patients with failed back surgery syndrome. *Clin Neurophysiol.* 2020;131(11):2578–2587. doi:10.1016/j.clinph.2020. 07.024
- 68. De Andres J, Monsalve-Dolz V, Fabregat-Cid G, et al. Prospective, randomized blind effect-on-outcome study of conventional vs high-frequency spinal cord stimulation in patients with pain and disability due to failed back surgery Syndrome. *Pain Med.* 2017;18 (12):2401–2421. doi:10.1093/pm/pnx241
- 69. Al-Kaisy A, Palmisani S, Smith TE, et al. Long-term improvements in chronic axial low back pain patients without previous spinal surgery: a cohort analysis of 10-kHz high-frequency spinal cord stimulation over 36 months. *Pain Med.* 2018;19(6):1219–1226. doi:10.1093/pm/pnx237
- 70. Fishman MA, Calodney A, Kim P, et al. Prospective, multicenter feasibility study to evaluate differential target multiplexed spinal cord stimulation programming in subjects with chronic intractable back pain with or without leg Pain. *Pain Pract.* 2020;20(7):761–768. doi:10.1111/papr.12908
- 71. Tilley DM, Lietz CB, Cedeno DL, Kelley CA, Li L, Vallejo R. Proteomic modulation in the dorsal spinal cord following spinal cord stimulation therapy in an in neuropathic pain model. *Neuromodulation*. 2020. doi:10.1111/ner.13103
- Ruiz-Sauri A, Orduña-Valls JM, Blasco-Serra A, et al. Glia to neuron ratio in the posterior aspect of the human spinal cord at thoracic segments relevant to spinal cord stimulation. J Anat. 2019;235 (5):997–1006. doi:10.1111/joa.13061
- 73. Vallejo R, Tilley DM, Vogel L, Benyamin R. The role of glia and the immune system in the development and maintenance of neuropathic pain. *Pain Pract.* 2010;10(3):167–184. doi:10.1111/j.1533-2500.20 10.00367.x
- 74. Chapman KB, Groenen PS, Vissers KC, van Helmond N, Stanton-Hicks MD. The pathways and processes underlying spinal transmission of low back pain: observations from dorsal root ganglion stimulation treatment. *Neuromodulation*. 2020. doi:10.1111/ ner.13150
- 75. Chapman KB, Groenen PS, Patel KV, Vissers KC, van Helmond N. T12 dorsal root ganglion stimulation to treat chronic low back pain: a case series. *Neuromodulation*. 2020;23(2):203–212. doi:10.1111/ ner.13047
- 76. Kallewaard JW, Edelbroek C, Terheggen M, Raza A, Geurts JW. A prospective study of dorsal root ganglion stimulation for non-operated discogenic low back pain. *Neuromodulation*. 2020;23 (2):196–202. doi:10.1111/ner.12937
- 77. Huygen F, Liem L, Cusack W, Kramer J. Stimulation of the L2-L3 dorsal root ganglia induces effective pain relief in the low back. *Pain Pract.* 2018;18(2):205–213. doi:10.1111/papr.12591
- Weiner RL, Yeung A, Montes Garcia C, Tyler Perryman L, Speck B. Treatment of FBSS low back pain with a novel percutaneous drg wireless stimulator: pilot and feasibility Study. *Pain Med.* 2016;17 (10):1911–1916. doi:10.1093/pm/pnw075
- 79. Sivanesan E, Bicket MC, Cohen SP. Retrospective analysis of complications associated with dorsal root ganglion stimulation for pain relief in the FDA MAUDE database. *Reg Anesth Pain Med.* 2019;44 (1):100–106. doi:10.1136/rapm-2018-000007
- Deer T, Pope J, Hunter C, et al. Safety analysis of dorsal root ganglion stimulation in the treatment of chronic pain. *Neuromodulation*. 2020;23(2):239–244. doi:10.1111/ner.12941

- Deckers K, De Smedt K, Mitchell B, et al. New Therapy for refractory chronic mechanical low back pain-restorative neurostimulation to activate the lumbar multifidus: one year results of a prospective multicenter clinical trial. *Neuromodulation*. 2018;21(1):48–55. doi:10.1111/ner.12741
- Ranger TA, Cicuttini FM, Jensen TS, et al. Are the size and composition of the paraspinal muscles associated with low back pain? A systematic review. *Spine J.* 2017;17(11):1729–1748. doi:10.1016/j. spinee.2017.07.002
- 83. Sions JM, Crippen DC, Hicks GE, Alroumi AM, Manal TJ, Pohlig RT. Exploring neuromuscular electrical stimulation intensity effects on multifidus muscle activity in adults with chronic low back pain: an ultrasound Imaging–Informed Investigation. *Clinical Medicine Insights: Arthritis and Musculoskeletal Disorders*. 2019;12:1179544119849570. doi:10.1177/1179544119849570
- 84. Cohen S, Gilmore C, Kapural L, et al. Percutaneous peripheral nerve stimulation for pain reduction and improvements in functional outcomes in chronic low back pain. *Mil Med.* 2019;184(Suppl Supplement 1):537–541. doi:10.1093/milmed/usy310

Journal of Pain Research

Publish your work in this journal

The Journal of Pain Research is an international, peer reviewed, open access, online journal that welcomes laboratory and clinical findings in the fields of pain research and the prevention and management of pain. Original research, reviews, symposium reports, hypothesis formation and commentaries are all considered for publication. The manuscript

management system is completely online and includes a very quick and fair peer-review system, which is all easy to use. Visit http:// www.dovepress.com/testimonials.php to read real quotes from published authors.

Submit your manuscript here: https://www.dovepress.com/journal-of-pain-research-journal

Dovepress