Cervical Spinal Cord and Dorsal Nerve Root Stimulation for Neuropathic Upper Limb Pain

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ABSTRACT: Background: Spinal cord stimulation (SCS) is a well-established treatment for chronic neuropathic pain in the lower limbs. Upper limb pain comprises a significant proportion of neuropathic pain patients, but is often difficult to target specifically and consistently with paresthesias. We hypothesized that the use of dorsal nerve root stimulation (DNRS), as an option along with SCS, would help us better relieve pain in these patients. Methods: All 35 patients trialed with spinal stimulation for upper limb pain between July 1, 2011, and October 31, 2013, were included. We performed permanent implantation in 23/35 patients based on a visual analogue scale pain score decrease of ≥50% during trial stimulation. Results: Both the SCS and DNRS groups had significant improvements in average visual analogue scale pain scores at 12 months compared with baseline, and the majority of patients in both groups obtained ≥50% pain relief. The majority of patients in both groups were able to reduce their opioid use, and on average had improvements in Short Form-36 quality of life scores. Complication rates did not differ significantly between the two groups. Conclusions: Treatment with SCS or DNRS provides meaningful long-term relief of chronic neuropathic pain in the upper limbs.

Keywords: Neurosurgery, Pain, Spinal Cord, Chronic Pain, Pain Control, Spinal Cord Stimulation, Neuromodulation, Nerve Root Stimulation
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Spinal cord stimulation (SCS) is a well-established treatment for chronic refractory neuropathic pain. There is a large body of evidence supporting its use, including several high-quality randomized controlled trials, mainly regarding stimulation of the thoracic spinal cord to treat lower limb pain resulting from failed back surgery syndrome.\textsuperscript{1,2} and complex regional pain syndrome (CRPS).\textsuperscript{3,4} Some areas of the body are thought to be more difficult than others to cover consistently and specifically with paresthesias, such as the upper limbs, groin, and low back.\textsuperscript{5,6} Given that upper limb pain comprises a significant proportion of neuropathic pain patients,\textsuperscript{7} this is a population that has historically been undertreated with neuromodulation, and further study would likely result in significant improvements to their care.

Two important issues have been discussed in the literature regarding the placement of an electrode and stimulation of the cervical spinal cord. First, the spinal cord enlargement from C3 to C7, coupled with this region’s susceptibility to degenerative changes and narrowing of the spinal canal, results in a shallow surrounding cerebrospinal space.\textsuperscript{5} This has raised concerns that...
electrode placement in this region has a potentially increased risk of neurologic injury and other complications, compared to electrodes in the thoracic and lumbar spine. Second, variation of stimulation amplitude and coverage with body position changes is a common patient complaint with spinal stimulation. This issue is potentially exacerbated by the mobility of the cervical spine and can require awkward postures to maintain consistent stimulation, limiting the overall effectiveness of the device.

Prior studies have reported success in treating upper limb CRPS and other forms of neuropathic pain with spinal cord stimulators. In our clinical practice, we have found that spinal dorsal nerve root stimulation (DNRS) is an effective technique for providing focused and consistent coverage for pain distributions and locations that are less amenable to treatment by traditional SCS. This technique uses the same stimulating electrodes and is easily converted to and from SCS during intraoperative testing, therefore it provides the surgeon with increased flexibility for targeting the pain area. In this paper, we report our experience with 35 patients treated for upper limb pain between July 1, 2011, to October 31, 2013, with a new trial of either SCS or DNRS.

METHODS

The study involved a single-center, open-label design and was approved by the Research Ethics Board at Western University, London, Canada. All patients who had been treated at University Hospital, London between July 1, 2011, and October 31, 2013, with a new implantation of a spinal stimulation trial system for upper limb pain entered the study.

Patient Selection

Patients were 18 years of age or older and had been referred for surgical management of upper limb pain that failed to respond to conservative measures including medication, psychological therapy, physical therapy, nerve blocks, and/or pain management programs. Exclusion criteria included pain in a distribution beyond the upper limb; another clinically significant or disabling chronic pain condition; an expected inability to manage or operate the SCS system; a history of a coagulation disorder; evidence of CRPS and other forms of neuropathic pain with spinal cord stimulators. Prior studies have reported success in treating upper limb CRPS and other forms of neuropathic pain with spinal cord stimulators. In our clinical practice, we have found that spinal dorsal nerve root stimulation (DNRS) is an effective technique for providing focused and consistent coverage for pain distributions and locations that are less amenable to treatment by traditional SCS. This technique uses the same stimulating electrodes and is easily converted to and from SCS during intraoperative testing, therefore it provides the surgeon with increased flexibility for targeting the pain area. In this paper, we report our experience with 35 patients treated for upper limb pain between July 1, 2011, to October 31, 2013, with a new trial of either SCS or DNRS.

Patient Selection

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Procedures

Treatment involved a trial period of 3 weeks, during which the programming of the device was adjusted and the patient’s response to the therapy was monitored. The trial electrodes were then removed and there was a mandatory washout period of at least two weeks. Permanent implantation followed successful trials, defined as a visual analogue scale (VAS) decrease ≥50% or a sufficiently large pain reduction to have a significant effect on the patient’s quality of life. The stimulation technique was chosen was based on the geographic distribution of the pain as well as intraoperative test stimulation results. In general, for the upper limb, nerve root stimulation was used in patients where the pain area was limited to one to four dermatomes. Electrodes were anchored to the thoracolumbar fascia using the standard technique with Boston Scientific’s Clik anchors. Examples of the placement of SCS and DNRS electrodes in the cervical spine are shown in Figures 1 and 2, respectively.

Data Collection and Analysis

Average pain intensity was assessed on a 10 cm VAS in which the point furthest to the left represented no pain (score: 0) and that furthest to the right represented the worst possible pain (score: 10). Quality of life was assessed using the Short Form (SF)-36 questionnaire. These endpoints were assessed prospectively at the following time points: before trial stimulation, three weekly visits during the trial, before permanent implantation, at 3-, 6-, and 12-month follow-up visits, and annually thereafter. Evaluations were performed by the pain specialist neurosurgeon, a nurse trained in neuromodulation, or a research student. Results are reported at 12 months, with the exception of average VAS scores, which are plotted at 3-, 6-, and 12-month follow-up. Average values are presented as ± standard error of the mean.

RESULTS

Patient Population and Trial Results

There were a total of 35 eligible patients, of which 18 had trials for DNRS, 15 for SCS, and 2 for both DNRS and SCS. Three patients had multiple trials, two of which were to attempt different stimulation techniques, whereas one patient had to have his first trial operation aborted because of discomfort. Twenty-three patients (66%) went on to permanent implantation, of which 11 received DNRS, 11 SCS, and one both.

Table 1 shows a breakdown of baseline characteristics for the SCS and DNRS groups of patients (the patients who received both were excluded from analysis because of small sample size). The two groups did not differ significantly in baseline VAS pain level, morphine equivalent daily dose (MEDD), SF-36 quality of life, employment status, age at implant, or time since pain onset. There was a greater proportion of female patients in the SCS group, of which 67% were female, compared with 44% of the DNRS patients. The most common pain diagnosis in each group was CRPS, followed by neuropathic pain not otherwise specified. The latter diagnosis included patients with posttraumatic, postsurgical, and spontaneous pain. There were two SCS patients with central neuropathic pain (poststroke and postspinal cord injury), and one DNRS patient with postherpetic neuralgia. All patients received electrodes placed in the cervical spine, and one DNRS patient also had an additional electrode in the thoracic spine (for both the trial and permanent implants).

Pain Reduction

Average VAS pain scores over 12 months of follow-up for each group of patients who received permanent implants are shown in Figure 3. At baseline, the average VAS was 7.8 (±1.2) for the SCS group and 7.6 (±1.2) for the DNRS group. At 12 months, the averages score had decreased to 2.7 (±0.6) for the SCS patients and 3.6 (±1.6) for the DNRS patients. At each of 3, 6, and 12 months, significant improvements from baseline pain levels were seen for both groups, and there were no statistically significant differences between the two groups.

Patients were categorized as responders, partial responders, and nonresponders based on pain reduction of ≥50%, 30% to 50%, and <30%, respectively. At 12 months, each patient group had pain data for nine patients, which is shown in Figure 4. In the
SCS group, six of nine (67%) were responders, two (22%) partial responders, and one (11%) was a nonresponder. In the DNRS group, six of nine were responders (67%) and three (33%) were partial responders.

Secondary Endpoints

At baseline, both patient groups had large impairments in all components of the SF-36 questionnaire for health-related quality of life, without significant differences between the two groups. There were five SCS patients and four DNRS patients who had SF-36 scores available at both baseline and 12-month follow-up. Figure 5 shows the difference between 12-month scores and baseline scores for each group. Both patient groups had improvements in the average score for all nine components, with the DNRS patients generally having larger improvements. Because of the small sample size, confidence intervals were not included in the figure. The only score change that reached statistical significance was the physical functioning component for the SCS group.

Both groups of patients were taking large doses of opioids at baseline, with an average MEDD of 218 mg (±251) for SCS patients and 170 mg (±284) for DNRS patients. At 12 months, the average MEDD had decreased to 98 mg (±126) and 144 mg (±259) for each group, respectively. Figure 6 shows the opioid doses at 12 months compared with baseline, divided into patients who had decreased, increased, and had no change in their dose, as well as patients who were never taking opioids. Of 10 SCS patients with data, five had decreased their MEDD, three had increased, and two were never on opioids. Of nine DNRS patients, six had decreased their MEDD and the remaining three were never on opioids.

At 12 months, five of six DNRS patients responded yes when asked if they were “satisfied with SCS treatment” and six of six
responded yes to “recommend to a friend?” Among SCS patients, seven of seven responded yes to both questions. On a clinical global improvement scale on which 1 represented “very much improved” and 7 “very much worse,” the average score was 1.75 (±0.71) for SCS patients and 1.67 (±1.03) for DNRS patients.

Complications and Revision Operations

During the trial period, the most common complications were superficial skin infection and lead migration, as shown in table 2. Infection occurred in two of 15 SCS patients (13%) and three of 18 DNRS patients (17%). One of these patients required surgical management of the infection, whereas the other four were treated just with oral antibiotics. Lead migration occurred in three of 15 (20%) SCS patients and one of 18 (6%) DNRS patients. Additionally, one SCS patient had to have a trial implant operation aborted because of discomfort, and one DNRS patient had a dural puncture during permanent implant, with resulting cerebrospinal leak and headache. During the permanent implant follow-up, there were a total of four revision operations for SCS patients and seven for DNRS patients. Three patients required multiple revision operations—one SCS patient had two revisions, one DNRS patient had two, and one DNRS patient had four. One SCS patient had his device removed because of lack of use. One DNRS patient had his electrode converted to an SCS system (he continued to be analyzed in the DNRS group). Otherwise, the reasons for revision were lead migration, implantable pulse generator malfunction, or attempted improvement of coverage.

DISCUSSION

Our study is the first to evaluate patients treated with dorsal nerve root stimulation of the cervical spine for upper limb pain. At 12 months, both the SCS and DNRS groups of patients had significant reductions in average VAS pain scores and the majority of patients obtained ≥50% pain reduction. In both groups, the majority of patients were able to reduce their opioid dose, and on average they had improvements in all subscores of the SF-36 questionnaire for quality of life. On patient satisfaction surveys, most patients indicated that they were satisfied with treatment and...
would recommend it to a friend; the average clinical global impression was between “much improved” and “very much improved.”

The technique for both SCS and DNRS at our centre uses the same hardware and stimulating electrodes. We must emphasize the need to place the electrode as far lateral as possible within the spinal canal for DNRS. Also, we take great care to ensure that the DNRS electrodes occupy the dorsal half of the neural foramina before stimulation to avoid ventral motor rootlet stimulation. Avoiding stimulation of the motor root also requires unique programming parameters. For dorsal root stimulation, we use a pulse width of 80 to 90 ms, compared with 250 to 700 in the SCS patients. Amplitude for DNRS is generally less than 2 mA, whereas it is highly variable for SCS.

Early in our experience, SCS was the initially trialed paradigm. Given current electrode technology, we can easily cover four cervical dermatomes/roots with DNRS. Therefore, if the pain area is confined to the C5-C8 dermatomes, we can cover the entire upper limb with DNRS. We have found DNRS to be much less positional overall and for upper limb pain it has become our default technique where possible. Although we did not specifically collect data regarding positional variations in stimulation amplitude and

### Table 1: Baseline patient characteristics

<table>
<thead>
<tr>
<th></th>
<th>SCS N = 15</th>
<th>95% CI</th>
<th>DNRS N = 18</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age in years: mean, SD</td>
<td>44.0</td>
<td>9.7</td>
<td>5.4</td>
<td>46.5</td>
</tr>
<tr>
<td>Years since pain onset: mean, SD</td>
<td>7.1</td>
<td>4.2</td>
<td>2.3</td>
<td>5.3</td>
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<td>Sex female: n, %</td>
<td>10</td>
<td>67</td>
<td>8</td>
<td>44</td>
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<tr>
<td>Currently employed: %</td>
<td>43</td>
<td></td>
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</tr>
<tr>
<td>VAS: mean, SD</td>
<td>7.8</td>
<td>1.2</td>
<td>0.7</td>
<td>7.6</td>
</tr>
<tr>
<td>MEDD: mean, SD</td>
<td>218</td>
<td>251</td>
<td>139</td>
<td>170</td>
</tr>
<tr>
<td>Diagnosis: n, %</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>CRPS</td>
<td>9</td>
<td>60</td>
<td>9</td>
<td>50</td>
</tr>
<tr>
<td>Neuropathic not otherwise specified</td>
<td>4</td>
<td>27</td>
<td>8</td>
<td>44</td>
</tr>
<tr>
<td>Central neuropathic</td>
<td>2</td>
<td>13</td>
<td>0</td>
<td>0</td>
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<tr>
<td>Postherpetic neuralgia</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>6</td>
</tr>
<tr>
<td>SF-36: mean, SD</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Physical functioning</td>
<td>58</td>
<td>16</td>
<td>15</td>
<td>49</td>
</tr>
<tr>
<td>Role-physical</td>
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<td>9</td>
<td>9</td>
<td>16</td>
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<tr>
<td>Bodily pain</td>
<td>19</td>
<td>21</td>
<td>19</td>
<td>18</td>
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<tr>
<td>General health</td>
<td>70</td>
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<td>48</td>
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<tr>
<td>Vitality</td>
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<td>Social functioning</td>
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<tr>
<td>Role-emotional</td>
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<td>24</td>
<td>12</td>
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<tr>
<td>Mental health</td>
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<td>17</td>
<td>41</td>
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<tr>
<td>Health transition</td>
<td>43</td>
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<td>22</td>
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</tr>
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</table>

CI = confidence interval; SD = standard deviation.
effectiveness, anecdotally, when both techniques are tested in individual patients, they find DNRS much less variable during typical neck movements. Both surgeons have had the same experience and agree on this. Patients with upper limb pain radiating into the neck or scapula are generally trialed with SCS first.

The most common complications were lead migration and superficial skin infection (during the trial period with exposed leads). Taking into account the relatively small sample size in this article and potential for random variation in numbers, our complication rate did not vary significantly between the two groups of patients and was consistent with our rate for patients with stimulators at other spine levels as well as reported complication rates in the literature. There were more overall revision operations in the DNRS patient group; however, this figure is skewed by the fact that one patient required four revisions from repeated lead fractures and migration. During the trial period we had infections in five patients (one requiring surgical management) of 35 patients who received a total of 40 trial operations. Although our trial period of 3 weeks is longer than what is done at other centres and can increase the risk of wound infection, we feel that a longer trial has value. We have found that there are some other centres and can increase the risk of wound infection, and this study found no significant difference in such a small sample rate. That said, it is possible that the electrode is better anchored by the shape of the lateral epidural space and it does subjectively feel more secure during insertion.

Our study has a number of strengths. It includes a large number of consecutive patients in a clinical setting, with a broad range of pain diagnoses, compared with other studies that have a more homogenous patient sample. Our patients were referred by family doctors or pain specialists and had severe long-term pain that had failed medical therapy and other forms of treatment. We had no unusual exclusion criteria for upper limb patients other than our standard clinical criteria for appropriate neuromodulation patients (listed in the Methods section). For analysis, we have separated the patients with DNRS and traditional SCS. Our results indicate that, contrary to previous assertions that nerve root stimulation is not effective, this lesser known technique performs equally well in appropriately selected patients.

The main limitation is that this is a self-funded study with limited resources; therefore, not all data are obtained at every clinic visit. This was particularly apparent for the SF-36 questionnaire; of 22 patients receiving permanent implants, only nine had SF-36 data at both baseline and at 12 months, whereas 19 had VAS data at this time point. It would have been preferable to have an independent third party doing patient evaluations, but this was not feasible given our resources. Our patients were not randomized to treatment groups but were assigned based on which technique was expected to be more effective for the individual based on pain distribution and intraoperative testing.

Prior studies have been published that have obtained similar results in pain reduction. A retrospective study of 23 patients with cervical spinal cord stimulation published in 2012 found that average pain scores decrease from 6.8 to 2.8, and a 2003 prospective study of 41 patients with either neuropathic or ischemic pain reported that 68% of patients obtain >50% pain relief. A prospective study published in 2004 by Forouzanfar et al. evaluated 36 CRPS patients, 19 of whom were treated with electrodes placed in the cervical spine and 17 in the lumbar spine. As with our findings comparing our upper limb patients with those treated with stimulators in other locations, this study found no significant difference in either pain relief or complication rate between the two groups.
An excellent paper was published by Deer et al. in 2014, outlining their prospective trial of 38 patients receiving cervical spinal cord stimulation, as well as a systematic literature review on the topic. Twelve studies were found that fulfilled their inclusion criteria, including a total of 221 patients. This included four prospective trials, four retrospective studies, and four case series. The most commonly cited adverse events were hardware malfunction (17.8%), lead migration (13.9%), and lead breakage (6.7%). Despite the fears that the cervical spine could have greater risk of cord compression during implantation, there were no reports of neurologic damage in the 180 patients for whom adverse events were reported. This review found no significant difference in the complication rate for cervical electrodes compared with previously published rates for thoracic and lumbar electrodes.

In summary, these findings indicate that SCS and DNRS in the cervical spine can provide good pain relief in upper limb neuropathic pain, as well as enable opioid use reduction and improve health-related quality of life. Despite concerns that electrodes in the cervical spine could be more predisposed to complications than those at other levels, our complication rates in this study are consistent with those reported in the literature for both stimulation at the cervical spine and at other spinal levels. Most important, no patient developed a new neurological deficit during the study.

DISCLOSURES

The authors have nothing to disclose.

STATEMENT OF AUTHORSHIP

AL collected and entered data into the database, interpreted and analysed data, conducted a literature review, and wrote the manuscript. AP participated in development of the study and helped to evaluate and edit the manuscript. KM acted as corresponding author, supervised the study, interpreted data, and edited the manuscript.

REFERENCES