Effect of Spinal Cord Stimulation for Patients with Post-Zoster Neuralgia on the Trunk: A Case Report

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Post-zoster neuralgia is an intractable pain by medical treatment. However, practice guidelines for invasive procedures of medically intractable post-zoster neuralgia are unclear, and it is not clear to apply to individual cases. In our study, the effect of spinal cord stimulation (SCS) for patients with post-zoster neuralgia was described in old patients with episodic pain with constant pain, which have been poorly controlled with oral and invasive treatments. We used an SCS device (Synergy2 IPG, model 7427; Medtronic Inc, Minneapolis, MN) equipped with an electrode (1×8 standard Lead model 977A290; Medtronic Inc, Minneapolis, MN). One lead was placed in the left lower margin of the T12 vertebral body (pulse width 60 ms, pulse rate 20 Hz, amplitude 2.5 V) and it covered on the distribution to left 10th, 11th, 12th thoracic dermatomes. VAS decreased to 3-4 after SCS compared with baseline VAS and was sustained by 3-4 for 1 week and by 3-4 at 1 month. After 6 months, the pain remained at 4-5. In morphine consumption, the baseline doses were 200 mg/day that significantly reduced to 120 mg/day and 90 mg/day after 1-month and 6-months post-procedure. The dose of gabapentin (1,800 mg/day) before treatment. The dose of gabapentin in the first month did not significantly decrease in taking gabapentin (1,200 mg/day) compared with the baseline dose (1,200 mg/day). Consumption after 6 months was also significantly reduced by 900 mg/day.

Key Words: Post-zoster neuralgia; Postherpetic neuralgia, Pulsed radiofrequency; Spinal cord stimulation

INTRODUCTION

A small number of patients experience pain during 4 weeks of herpes zoster infection, and post-zoster neuralgia occurs in 10% of the patient after herpes zoster 2). The combination therapy of pharmacologic agents is more effective than any monotherapy for the management of post-zoster neuralgia 4). However, invasive practical guidelines for medically intractable patients failing current drug therapy are unclear, and it is not clear to apply to the individual instructions 3). Therefore, the objective of this report is to evaluate the efficacy of spinal cord stimulation from a case with post-zoster neuralgia on left 10th, 11th, 12th thoracic dermatomes

CASE REPORT

A 68-year-old male had a history of healing in herpes zoster infection with rash and blister on left 10th, 11th, 12th thoracic dermatomes. This patient with severe left trunk pain for 126 months was referred to our pain clinic for adjustable pain management. Post-zoster neuralgia began following bullae and cluster in a herpes zoster infection on the distribution to left 10th, 11th, 12th thoracic dermatomes. He described the pain as itching and lancinating sensation with constant and severe (VAS 7 to 9 as being the most severe). The patient did not respond to various drug treatments and combination treatment for relief of trunk pain: ibrudon 60 mg bid, acetaminophen/hydrocodone 5/500 tid, pregabalin 150 mg bid, topical agents, the combination treatment of injection to scar and ICNB. For the relief of his post-zoster neuralgia, We used an SCS device (Synergy2 IPG, model 7427; Medtronic Inc, Minneapolis, MN) equipped with an electrode (1×8 standard Lead model 977A290; Medtronic Inc, Minneapolis, MN). One lead was placed in the left lower margin of the T12 vertebral body (pulse width 60 ms, pulse rate 20 Hz, amplitude 2.5 V), and is covered on the distribution to left 10th, 11th, 12th thoracic dermatomes (Figs. 1, 2). After the trial stimulation, his visual analog score (VAS) decreased from 9 to 4. SCS-induced paresthesia covered his painful areas fully enough. The VAS of baseline status was 7-9 before SCS in the management of episodic pain with constant pain in patients with post-zoster neuralgia. VAS decreased to 3-4 after SCS compared with baseline VAS and was sustained by 3-4 for 1 week and by 3-4 at 1 month. After 6 months, the pain remained at 4-5. In morphine consumption, the baseline doses
Fig. 1. (A) The root space of thoracic spine was accessed at L2-3 interspace and the C-arm rotated until the rootlet space was easily identified. (B) A 14-gauge, 10-cm Tuohy needle was advanced under fluoroscopic guidance with a direct approach.

Fig. 2. An electrode (1×8 standard Lead model 977A290; Medtronic Inc, Minneapolis, MN) was placed in the left lower margin of the T12 vertebral body and an electrode (1×8 standard Lead model 977A290; Medtronic Inc, Minneapolis, MN) (A: D-spine AP view, B: D-spine lateral view).

were 200 mg/day that significantly reduced to 120 mg/day and 90 mg/day after one 1-month and 6-months post-procedure. The dose of gabapentin (1,800 mg/day) before treatment. The dose of gabapentin in the first month did not significantly decrease in taking gabapentin (1,200 mg/day) compared with the baseline dose (1,200 mg/day). Consumption after 6 months was also significantly reduced by 900 mg/day.

**DISCUSSION**

During the acute herpes zoster infection, damage of peripheral nerves lowered threshold for pain activation from nociceptive organ and spontaneously ectopic discharges by destructed nerve cause peripheral and central sensitization\(^5\). Neurons of dorsal horn in spinal cord are also altered by response to the activity and inflammatory changes in peripheral nerves impaired descending inhibition of pain sensation and led to central sensitization\(^6,10\).

Treatment guidelines of post-zoster neuralgia suggested that medical treatment in post-zoster neuralgia is excellent in an effect of combination therapy of pharmacologic agents rather than a single therapy\(^3\). Looking at the drug treatment guidelines for post-zoster neuralgia issued by the International Association for the Study of Pain (IASP), First-line treatments include TCAs (tricyclic antidepressants), pregabalin and gabapentin, and topical lidocaine, 5% opioid patch, capsaicin cream, and 8% capsaicin patch are additionally recommended. Traditionally, anticonvulsants as carbamazepine and valproic acid showed an effect in some patients, and botulinum toxin had been applied to pain management. SSRIs (Selective serotonin reuptake inhibitors) and SNRIs (serotonin-norepinephrine reuptake inhibitors) have been applied to the medical treatment of post-zoster neuralgia\(^9,11\).

In comparison, the subcutaneous injection of lidocaine and triamcinolone is a safe and straightforward method to block affected peripheral nerves and reduce inflammation. In the literature, the pain NRS scores decreased significantly in all patients with post-zoster neuralgia one month after rash onset. Still, patients who received standard treatment plus subcutaneous injection showed substantially more significant improvement in pain scores than those who received standard therapy alone\(^1\).

Some presented 2 subacute PHN cases within 3 months of the onset of herpes zoster, in which limited duration SCS was effective in alleviating the pain. Increasing drug dosages or switching drugs was difficult because both patients were elderly\(^8\).

The use of SCS within 0.5 to 2 months from the onset of herpes zoster improved VAS scores to zero, and the patients could complete the therapies in 2.5 months. Responses in post-zoster neuralgia patients were not uniform; however, as some patients still had pain after 29 months of the treatment\(^7\). Some proposed to utilize SCS for approximately 2 weeks without permanent implantation of the pulse generator. They showed that patients who were within 3 months from the onset of herpes zoster achieved pain relief\(^2\). Some reported that early intervention with SCS provided better pain relief as well as longer periods of effective control. Also, long-term pain alleviation and patient satisfaction were superior when the SCS therapies were started earlier\(^9\). In another report, the cost-effectiveness of the SCS was not found to be inferior compared to drug therapies\(^13\).

**CONCLUSION**

In summary, the efficacy of spinal cord stimulation from
post-zoster neuralgia represents an effective and safe method for post-zoster neuralgia on the spinal nerve. The benefits of SCS include minimal invasiveness that provides rapid relief of pain and improves quality of life.

REFERENCES