The effect of cervical spinal cord stimulation on cervical spinal nerve root/brachial plexus injury

Servikal spinal kord stimulasyonunun servikal spinal sinir kökü/brakial pleksus yaralanmasını üzerine etkisi

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Summary
Spinal cord stimulation (SCS), has been used for the treatment of chronic pain for almost 50 years. There have been several reports regarding to thoracic and lumbar SCS, while fewer publications have been documented for cervical SCS administration. In this article, we presented patient satisfaction after cervical SCS application in patients with chronic upper extremity pain. Three patients with cervical spinal nerve root or brachial plexus injury who reported no pain relief with previous treatment modalities were identified. The patients were performed percutaneous cervical spinal cord stimulation. Cervical SCS implantation was successful in the first and the second patients, and with respect to decrease in VAS scores, analgesic drug requirements, and an increase in their quality of life while the third patient had no pain relief. Cervical spinal cord stimulation may be an effective treatment option in upper extremity chronic pain syndromes as in lower extremity pain syndromes treated with thoracolumbar SCS. Further studies will guide the effectiveness and applicability of cervical SCS in the future.

Keywords: Brachial plexus injury; cervical nerve root injury; chronic pain; neuromodulation; neuropathic pain; spinal cord stimulation

Introduction
Spinal cord stimulation (SCS) was first performed by Shealy et al in 1967 and it has been used for the treatment of chronic pain for almost 50 years.[1,2] The technique became widespread in 1975 with the development of percutaneous electrodes.[1] Modalities such as surgical ablative operations, pharmacotherapy and physical therapy are applied in the treatment of chronic pain but may be insufficient. Spinal cord stimulation produce neuromodulatory effect by stimulating spinal cord with low intensity electrical impulses and can significantly change the perception of pain in patients with chronic pain.[3] There have been several long-term and successful reports of thoracic and lumbar spinal cord stimulation, while fewer publications have been documented for cervical SCS administration.[4] The aim of this case report is to evaluate and present patient satisfaction after cervical SCS application in patients with chronic upper extremity pain.

Case Report
Case 1 – A 36 year-old male patient, developed a C4 vertebral fracture and left carotid artery dissection as a result of a gunshot wound to the neck for four
years ago before admitting to pain clinic. The patient complained of aching, throbbing, neuropathic pain spreading to left arm and neck. His visual analog scale (VAS) was 8–9/10. On physical examination; severe pain, coldness, weakness found on his left shoulder, arm, forearm and hand, and there was hypoesthesia on the left C4, C5, C6 spinal nerve dermatome areas. Muscle strength on the left deltoid, triceps, flexor and extensor digits were 4/5. There was mild pain in the left leg, paresthesia in the right leg, temperature difference between the lower extremities, and he had impotence and urinary incontinence.

There were chronic, axonal degeneration involving the C5, C6 spinal nerve roots, left brachial plexus on electromyography. Cervical spinal MRI (2008) showed that C4-C5 facet segment fracture, left neural foramen obliteration was present. He was prescribed pregabaline 300 mg 2*1 p.o, SSRI (sertraline) 50 mg 1*1 p.o, baclofen 10 mg 1*2 p.o and oxycodone 10 mg 4*1 p.o before presented to clinic. He had a history of cervical epidural injection in an another clinic. But he had no pain relief despite all the treatments.

In April 2013, a SCS catheter was inserted into the epidural space at the level of T2-T3 thoracic vertebra and fixed in the epidural space at the C3-C4 cervical vertebra level, with the percutaneous electrode slightly to the left of the midline (Fig. 1). He felt paresthesia on the painful area and reported 70% pain relief during trial period, and permanent SCS implantation was placed four weeks later. Patient’s VAS score was 2–3/10 after permanent SCS implantation, and medical treatment was continued with pregabalin 300 mg 2*1 p.o and sertraline 50 mg 1*1 p.o. He reported no need for other analgesic drugs and improvement in his social life quality during long term follow-up.

**Case 2** – 58 year-old male patient admitted with a history of causalgia like, severe burning pain in his left shoulder and arm associated with traumatic cervical root injury, left brachial plexus lesion due to dislodgement of a metal part from a working machine in 1978, at the age of 18. His VAS score was 9/10. On his physical examination; loss of sensation in the left elbow, muscular atrophy in left arm, trophic changes in nails, dry skin, color changes, limitation of joint movements and severe pain, absence of deep tendon reflex, shoulder abductors and biceps muscle strengths were 2/5, deltoid, triceps, hand flexion-extension, and interosseous muscle strengths were 0/5 and flask paralysis were present.

There were complete conduction defect in left n. medianus, n. ulnaris, n. radialis, n. musculocutaneus, n. axillaris and left brachial plexus lesion on electromyography, nerve root expansion was seen in and around the left neural foramen at the C7-T1 and T1-T2 vertebral level in cervical computed tomography (CT), and in myelography, complete nerve root avulsion at the vertebral levels of C5-C6, C6-C7, C7-T1 and partial nerve root avulsion at the vertebral level of T1-T2 were observed. He was prescribed, fentanyl patch 100 mcg/48 hours tts, gabapentin 600 mg 2*1/2 p.o. In Germany, in 1979, he was performed deep brain stimulation, but the patient had had no benefit from this treatment. (MRI examination could not be performed due to this procedure.)

In 2004, SCS implantation was performed to the patient in our clinic whose pain was continued despite all the treatments. SCS catheter was inserted into the epidural space at the level of T2-T3 thoracic vertebrae and advanced to the C5-C6 cervical vertebra level and fixed in the same level of the epidural
space (Fig. 2). The patient reported VAS score 2 after the procedure and satisfied with the procedure by 70%. He received gabapentin 800 mg 2*1/2 p.o as medical treatment only, and he reported increase in his quality of sleep and quality of life.

In 2009 and 2017, due to completion of the generators life time, neurostimulatory pulse generator revision was also performed.

**Case 3** – A 46-year-old male patient was admitted with a complaint of left shoulder, arm and hand pain with burning, tingling, aching like pain after left brachial plexus injury due to traffic accident for 10 years ago.

Atrophic appearance was seen on his left shoulder, arm, forearm and sensory and motor deficit were present on his physical examination. Transcutaneous electrical nerve stimulation (TENS) was applied previously and gabapentin 600 mg 3*1 p.o, escitalopram (SSRI) 20 mg 1*1 p.o, tramadol 100 mg 2*1 p.o was prescribed as the analgesic treatment.

Left stellate ganglion block was performed three times at the 1.5 year, 4.5 years and 5.5 years after the brachial plexus injury. He had 40% pain relief from the first block but had no relief from subsequent procedures. Despite medical treatment and invasive procedures, it was decided to perform cervical SCS to the patient whose pain was ongoing as VAS 7/10.

EMG in May 2018; lesion at the level of brachial plexus with left cervical roots, involvement on proximal to dorsal root ganglion (avulsion?), partial degeneration in upper and lower truncus, complete axonal degeneration in peripheral nerves were present.

In cervicothoracic MRI; there were no other pathology other than protrusion and bulging in two levels of intervertebral disc. There were no pathology in dorsal spinal MRI and no contrast enhancement was observed.

In June 2018, SCS electrode was inserted into the epidural space at the level of T3-T4 thoracic vertebrae and fixed at the C4,5,6,7 (upper end of C7) cervical vertebra level of the epidural space (Fig. 3). During the cervical SCS trial period, the patient felt paresthesia in the painful area but he reported less than 50% pain relief. Different stimulation parameters applied for three weeks. Although the patient experienced paresthesia in painful areas during the trial period SCS electrode was removed, due to less than 50% pain relief. The patient was discharged with medical treatment and recommendation of continuation of physical therapy.

**Discussion**

Spinal cord stimulation (SCS) has become a preferred and increasingly widespread treatment modality in patients with chronic pain who have received various treatments previously. The basic theory of SCS is based on Melzack and Wall’s ‘gate control theory’; SCS-induced analgesia is multifactorial and has not
yet been fully elucidated. Despite there are many studies and case series in thoracolumbar SCS implantation, the literature is limited in cervical SCS implantation. It is thought that the narrowness of the epidural space in the cervical region, the higher risk of cord compression, hypermobility of the cervical spine may contribute to the limitation. In some respects, cervical SCS has some differences from the thoracolumbar SCS. First, the mobility of the cervical spine is higher and paresthesia changes are more likely to occur, but there is no large-scale study. Secondly, paresthesia of the trunk and lower extremities is possible due to the different anatomy of the dorsal column. Vallejo et al, in a series of 5 patients, reported generalized paresthesia in 3 patients. However, since these patients have pain in the lower extremities, paresthesia is seen as an additional benefit rather than an undesirable condition. Similarly, lower extremity pain was present in the first one of our cases and our patient reported a decrease in lower extremity pain. The absence of lower extremity stimulation in all patients who were performed cervical SCS can be explained based on the study by Freirabend et al. In their study on dorsal column morphometry, they concluded that fiber density increased from medial to lateral. Thus, they stated that the lateral fibers will be affected in the low stimulation intensity and the stimulation will be detected in the upper extremity, and the high stimulation intensity will affect the medial fibers and the stimulation will be detected in the trunk and lower extremity.

In a prospective study of Deer et al, about the safety and efficacy of cervical SCS, they stated that the mean pain palliation of 38 patients was more than 50% (66.8%) in one year after SCS implantation. So they proposed that cervical SCS is safe and effective. Similarly, in the study of Chivikula et al in a large number of (121) patients, 75% of the cervical SCS electrode was implanted permanently and the patients’ satisfaction rate was 72.4%. The rate of pain palliation was 58.7% about the cervical SCS implantation, like thoracic and lumbar areas. Kumar et al. evaluated 10 patients who were performed cervical SCS implantation, by the visual analog scale (VAS), Oswelt disability index (ODI) and Beck depression inventory (BDI) of the patients. They observed a significant improvement in VAS scores of 34%, 9.2% for ODI scores, and 29.6% for BDI scores for the long-term follow-up for about 7 years. And it is thought that the early application of SCS, including cervical SCS, in long-term chronic pain treatment will be more beneficial.

In addition, Garg A. and Danesh H. reported a case of cervical dorsal root ganglion (DRG) neuromodulation in a patient with complex regional pain syndrome who was previously performed cervical discectomy. During the procedure, they reported that the patient had scar tissue in the posterior epidural area due to the cervical discectomy operation previously, and incidentally, the lead tip was directed to the C6 foramen, so they fixed the electrode at the level of the C6 dorsal root ganglion. They reported 70% pain relief and stated that especially in distal extremity pain, stimulation of dorsal root ganglion may be an alternative treatment to spinal cord stimulation.

In our cases, cervical SCS implantation provided pain relief in the first and the second patients and an increase in their quality of life. The well-being of the second case has been going on for about 40 years and the termination of the intensive opioid requirement is considered highly valuable. Third patient had less than 50% pain relief and permanent implantation was not performed. Pain characteristics were similar in all three patients. The third patient’s inability to benefit from the procedure could not be attributed to the duration of pain because the duration of pain was quite different in all three patients, and 4, 40 and 10 years, respectively. However, the painful area, that including the neck, shoulder, arm, forearm and hand was quite wide, and the absence of muscle strength in these areas (0/5), lack of sensitivity in any way, absence of intact myelinated A-beta fibers and degeneration of the dorsal column fibers by the lesion in the proximal nerve roots, which is very close to the dorsal root ganglia, may have contributed to absence of pain relief.

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### Conclusion

Interventional neuromodulation methods are becoming increasingly widespread in chronic pain. Spinal cord stimulation has some advantages such as reducing the need for oral analgesics that have many side effects, reducing recurrent invasive pro-