Percutaneous Peripheral Nerve Stimulation for the Treatment of Chronic Pain Following Amputation


ABSTRACT Introduction: Chronic pain and reduced function are significant problems for Military Service members and Veterans following amputation. Peripheral nerve stimulation (PNS) is a promising therapy, but PNS systems have traditionally been limited by invasiveness and complications. Recently, a novel percutaneous PNS system was developed to reduce the risk of complications and enable delivery of stimulation without surgery. Materials and Methods: Percutaneous PNS was evaluated to determine if stimulation provides relief from residual and phantom limb pain following lower-extremity amputation. PNS leads were implanted percutaneously to deliver stimulation to the femoral and/or sciatic nerves. Patients received stimulation for up to 60 days followed by withdrawal of the leads. Results: A review of recent studies and clinical reports found that a majority of patients (18/24, 75%) reported substantial (≥50%) clinically relevant relief of chronic post-amputation pain following up to 60 days of percutaneous PNS. Reductions in pain were frequently associated with reductions in disability and pain interference. Conclusions: Percutaneous PNS can durably reduce pain, thereby enabling improvements in quality of life, function, and rehabilitation in individuals with residual or phantom limb pain following amputation. Percutaneous PNS may have additional benefit for Military Service members and Veterans with post-surgical or post-traumatic pain.

INTRODUCTION Post-amputation pain (PAP) is a substantial military health problem. Over 50,000 US Service members have been injured in military conflicts since 2001 (e.g., Operation Enduring Freedom [OEF], Operation Iraqi Freedom [OIF]), and as a result, over 1,600 major limb amputations have occurred.1,2 Due to advances in military medicine including improved body armor, faster access to advanced life support and theater-based medical facilities, and effective technological care in the field, the mortality rate of severely injured Service members is decreasing.3,4 As a result, there is an increasing prevalence of warriors surviving traumatic injuries and a strong emphasis on alleviating pain and improving quality of life in these individuals.5

Recent studies indicate that more than 85% of US Service members with combat-related traumatic amputations from OEF/OIF suffer from moderate to severe PAP.6,7 Pain is a leading cause of disability, and PAP in particular can be extremely debilitating. Only 3–18% of Veterans from Vietnam and OEF/OIF with amputations are able to perform high-impact aerobics.8 Post-amputation pain significantly decreases quality of life, increases the risk of depression, discouragement, and anger, and negatively affects interpersonal relationships and the ability to return to work (amputees have a low probability [2–7%] of returning to active duty).7,9–12 Pain also negatively affects a Service member’s ability to comfortably wear a prosthetic device. Over 40% of Vietnam and OIF/OEF Service members and Veterans with multiple lower limb loss suffer from pain while wearing their prosthetic devices, and some ultimately abandon their prosthetics, which negatively impacts their ability to perform physical activities and maximize their level of fitness.9 Poorly controlled pain is an important barrier that must be overcome in order to promote functional recovery and effective rehabilitative therapy.

Chronic PAP includes residual limb pain (RLP) and phantom limb pain (PLP). RLP can be caused by multiple sources after amputation, including tissue trauma associated with surgery, wound infection or poor healing, heterotopic ossification, and direct or referred mechanical pain from joint degeneration or prosthetic fit.13–15 In most cases, residual limb pain is nociceptive in nature. Although neuroplasticity can accompany any pain condition, PLP may be more commonly associated
with peripheral and central sensitization than RLP, including functional reorganization of nociceptive pathways in the spinal cord and brain, sensory remapping, expansion of receptive fields, and altered cortical representation of the limb.\textsuperscript{13-16} Pain may also be referred to the phantom limb from neuromas, radiculopathy, a proximal neural lesion, or even biomechanical non-neuropathic causes such as bursitis.\textsuperscript{14}

The various potential mechanisms underlying PAP were traditionally categorized as nociceptive pain (due to actual or threatened tissue damage) or neuropathic pain (due to a disease or lesion affecting the somatosensory nervous system) (Table I).\textsuperscript{13} For example, prosthogenic pain and heterotopic ossification are nociceptive in nature, whereas hyperalgesia and allodynia from neuroma growth are neuropathic mechanisms. Recently, the International Association for the Study of Pain introduced a third mechanistic category, nociplastic pain, that includes pain not otherwise related to tissue insult or a somatosensory neural lesion.\textsuperscript{17}

Some of the changes that are observed after amputation, such as maladaptive cortical plasticity, central sensitization, decreases in the activity of descending inhibitory systems, and abnormal activity in the sympathetic nervous system,\textsuperscript{16} may be considered nociplastic mechanisms that promote maladaptive pain in the absence of overt pathology and persist beyond tissue healing timelines (Table I). Table II illustrates differences in clinical characteristics among the three pain classifications.

### PERIPHERAL NERVE STIMULATION (PNS) IS A THERAPEUTIC OPTION FOR CHRONIC PAIN FOLLOWING AMPUTATION, BUT HAS TRADITIONALLY BEEN LIMITED BY COMPLICATIONS AND INVASIVENESS

Likely because of its complex, multifaceted nature, PAP has historically been a challenging condition to treat, and amputees often progress through a battery of management techniques and procedures without finding adequate relief. Many approaches have been used, including opioid and non-opioid oral analgesics, nerve blocks and radiofrequency therapies, spinal cord stimulation (SCS), and physical and psychological therapies, but clinical experience and controlled trials have not demonstrated consistent and effective pain management using these traditional methods.\textsuperscript{13,14,18-21} For example, opioid analgesics have shown some success in a few trials, but data are limited and many amputees either fail to achieve consistent long-term pain relief or suffer from side effects common to opioid medications, such as nausea, drowsiness, headache, insomnia, loss of libido, and addiction.\textsuperscript{18,21} Rising levels of opioid addiction and associated socioeconomic burdens have also prompted a reexamination of prescribing practices and highlight the need for non-opioid pain management options.

PNS has been used effectively for immediate and lasting relief of chronic pain, including in individuals with chronic pain following amputation.\textsuperscript{12-24} PNS has also been shown to improve sleep, increase quality of life and activity levels, allow a significant percentage (up to 50%) of patients to return to work, and reduce or eliminate dependence on opioid analgesia.\textsuperscript{25,26} In one review of 117 patients receiving PNS who were followed up to 53 months, 65% reported an increase in their activities of daily living and more than 75% were satisfied with therapy.\textsuperscript{27}

PNS can have a high success rate and positively impact function and quality of life through pain relief, but traditional methods of surgically placing the lead near or in direct contact with the nerve have historically been limited by invasiveness, complexity of surgical implantation, and risk of complications such as nerve damage, lead migration (24–33%), infection (1–5%), pain at the site of implantable pulse generator (IPG) (0.9–5.8%), and hardware or battery failure (1.6–2%).\textsuperscript{28} In one retrospective review of traditional PNS techniques, Ishizuka and colleagues found that 64% of patients required reoperation because initial pain relief following surgical implantation was lost due to lead migration, infection, or poor initial lead placement.\textsuperscript{29} Collectively, these complications contribute to an overall revision rate of 27% among patients undergoing surgical implantation of traditional PNS systems, nearly half of which (15%) require explantation.\textsuperscript{28}

### PERCUTANEOUS PNS IS DESIGNED TO ADDRESS MANY OF THE CHALLENGES PREVIOUSLY ASSOCIATED WITH PNS

It was theorized that a system designed specifically for the non-surgical, reversible implantation of leads remote from...
the target nerve in the peripheral nervous system would facilitate effective pain management while reducing or eliminating many of the common complications previously associated with PNS. In particular, analysis of stimulation thresholds suggested that implantation of leads remote from the target nerve (0.5–3 cm) would enable preferential activation of targeted (pain-relieving) fibers while avoiding activation of nontarget fibers that can cause discomfort and limit the utility or therapeutic window of PNS therapy.

An FDA-cleared percutaneous PNS system was developed with the intent of overcoming the challenges of lead migration and invasiveness that limited previous approaches to PNS by employing flexible, fine-wire, helically-coiled leads designed to resist migration (Fig. 1). Open-coil percutaneous leads have a history of over 40 years of effective use in the periphery and are designed to resist migration in part by promoting healthy tissue ingrowth between the coils to anchor the lead in tissue. The leads are pre-loaded within a needle-based introducer and percutaneously implanted, typically under ultrasound guidance. The leads are implanted remote from the target nerve (0.5–3 cm), minimizing the risk of nerve injury (Fig. 1). PNS therapy is delivered for up to 60 days, after which the leads are withdrawn.

**PERCUTANEOUS PNS REDUCES PAIN AND PAIN INTERFERENCE IN INDIVIDUALS WITH CHRONIC PAIN FOLLOWING AMPUTATION**

The feasibility of using percutaneous PNS for relieving chronic pain after amputation was first demonstrated by Rauck and colleagues in an NIH-funded case series study. Fourteen of 16 patients (88%) with RLP and/or PLP following lower-limb amputation reported immediate clinically-relevant pain relief following implantation of leads remote from the femoral and/or sciatic nerves. Leads were

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**TABLE II. Clinical Characteristics of Nociceptive, Neuropathic, and Nociplastic Post-Amputation Pain**

<table>
<thead>
<tr>
<th></th>
<th>Nociceptive Post-Amputation Pain</th>
<th>Neuropathic Post-Amputation Pain</th>
<th>Nociplastic Pain in Amputees</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Etiology</strong></td>
<td>Actual or potential tissue damage, referred pain from mechanical structures</td>
<td>Severing of nerve, neuroplastic changes in the peripheral and central nervous systems</td>
<td>Altered nociception despite no evidence of actual or threatened tissue damage, or evidence for a lesion affecting the somatosensory system. Trauma is a common antecedent to CRPS type I, uncommon for other types of nociplastic pain.</td>
</tr>
<tr>
<td><strong>Frequency</strong></td>
<td>Most common cause of residual limb pain</td>
<td>Most common cause of phantom limb pain</td>
<td>Infrequent stand-alone cause of post-amputation pain, though altered pain processing may accompany nociceptive and neuropathic postamputation pain</td>
</tr>
<tr>
<td><strong>Descriptors</strong></td>
<td>Throbbing, aching, pressure-like</td>
<td>Lancinating, shooting, electrical-like</td>
<td>Highly variable</td>
</tr>
<tr>
<td><strong>Sensory Changes</strong></td>
<td>Infrequent, outside of a nerve or nerve root distribution</td>
<td>Phantom sensations very common</td>
<td>Common, but often outside the distribution of nerve or tissue injury</td>
</tr>
<tr>
<td><strong>Hypersensitivity</strong></td>
<td>Uncommon except for hypersensitivity in the immediate area after trauma or amputation, often elicited by palpation of pain generator</td>
<td>Allodynia and hyperalgesia may be present in residual limb</td>
<td>Hallmark of the condition</td>
</tr>
<tr>
<td><strong>Location</strong></td>
<td>Proximal radiation frequent</td>
<td>Distal radiation common, telescoping often observed</td>
<td>Diffuse, outside the distribution of an injured nerve(s) or amputated body part</td>
</tr>
<tr>
<td><strong>Time course</strong></td>
<td>Acute postsurgical pain decreases over several weeks. Pain from other sources stabilizes or slightly diminishes over time, though referred pain from degenerative diseases may persist or worsen</td>
<td>Often experienced within 1 week of amputation, prevalence peaks within 2 years and remains stable or declines in intensity</td>
<td>Pain post-injury disproportionate to inciting event. Delays in diagnosis common.</td>
</tr>
<tr>
<td><strong>Paroxysms</strong></td>
<td>Exacerbations less common and often associated with specific activities (putting on prostheses, ambulation)</td>
<td>Exacerbations common and unpredictable</td>
<td>May be superimposed on low-grade continuous pain</td>
</tr>
<tr>
<td><strong>Autonomic signs</strong></td>
<td>Uncommon</td>
<td>Can occur in 1/3 to 1/2 of patients</td>
<td>Frequent in CRPS type I and other types of nociplastic pain</td>
</tr>
<tr>
<td><strong>Associated symptoms</strong></td>
<td>Psychiatric co-morbidities common</td>
<td>Psychiatric co-morbidities common</td>
<td>High co-prevalence rate of other nociplastic pain conditions. Cognitive deficits, psychiatric co-morbidities, fatigue, poor sleep and sensitivity to light and other stimuli common</td>
</tr>
</tbody>
</table>
implanted percutaneously under ultrasound guidance distal to the inguinal crease approximately 0.5–1 cm superficial or lateral to the femoral nerve (Fig. 1B), and transgluteally approximately 1–1.5 cm superficial or lateral to the sciatic nerve between the greater trochanter and ischial tuberosity. Nine patients continued use of the PNS system for two weeks and reported a 72% mean reduction of RLP and 81% mean reduction of PLP (Fig. 2).

Reductions in post-amputation pain were sustained following lead withdrawal throughout the 4-week follow-up period. In addition to pain relief, patients reported 81–83% mean reductions in RLP and 81% mean reduction of PLP (Fig. 2). Reductions in post-amputation pain were sustained following lead withdrawal throughout the 4-week follow-up period. In addition to pain relief, patients reported 81–83% mean reductions in RLP and 81% mean reduction of PLP (Fig. 2). Reductions in post-amputation pain were sustained following lead withdrawal throughout the 4-week follow-up period. In addition to pain relief, patients reported 81–83% mean reductions in RLP and 81% mean reduction of PLP (Fig. 2).

Following the initial feasibility study, a DoD-funded, multicenter, randomized, double-blind, placebo-controlled trial further evaluated percutaneous PNS for the treatment of chronic pain following amputation in 28 traumatic lower extremity amputees. Leads were placed 0.5–1 cm remote from the femoral nerve and 1–1.5 cm remote from the sciatic nerve using approaches similar to those described by Rauck et al. Gilmore and colleagues found that a significantly greater proportion of patients who received PNS for up to 60 days experienced substantial (≥ 50%) reductions in pain compared to patients who received placebo therapy (Fig. 2). These reductions in pain were achieved with concurrent reductions in opioid usage in some patients; the subset of subjects taking moderate-to-high opioid doses at baseline reported 71% average reductions in daily opioid consumption after up to 60 days of PNS. According to Gilmore et al., prospective follow-up in this study was ongoing and 5 patients had completed the 12-month follow-up at the time of writing (with statistical analysis pending completion of follow-up in additional patients). Substantial (≥ 50%) reductions in pain were sustained in 4 of 5 patients (80%) who had completed the 12-month follow-up visit to-date. Gilmore and colleagues also found significant improvements in function, as measured by pain interference, Patient Global Impression of Change (PGIC), and BDI-II. Patients
in the PNS therapy group reported average reductions in pain interference of \( \geq 4 \) points on a 0–10 rating scale (Fig. 2),\(^{39}\) which was more than four-times higher than the one-point threshold for a minimally important change.\(^{40}\) Pain interference was reduced by 75% compared to baseline, and 80% of patients reported \( \geq 50\% \) reductions. Patients receiving PNS also reported being “Much Improved” to “Very Much Improved” on the PGIC survey and had statistically significantly greater global improvement and greater decreases in depression as measured by BDI-II compared to patients receiving placebo.\(^{39}\) Disability and quality of life are significant issues for amputees including injured military Service members, and pain relief from percutaneous PNS may provide additional benefits by enabling greater function and return to activities of daily living previously affected by chronic pain after amputation.

In addition to studies that focused predominantly on individuals with traumatic amputations such as those that occur secondary to combat-related injuries, percutaneous PNS therapy has also been used to treat Veterans with non-traumatic amputations outside of the clinical research setting. The Hunter Holmes McGuire VA Medical Center (HHMVAMC) is one of seven Regional Amputation Centers in the Veteran’s Health Administration Amputation System of Care (ASoC).\(^{41}\) The ASoC was implemented in 2008 to help provide Veterans with limb loss with state-of-the-art care delivered with an interdisciplinary focus on pain management, residual limb care, and other aspects of rehabilitation.\(^{41}\) Three Veterans at HHMVAMC who experienced chronic pain following dysvascular or infection-related lower extremity amputation have undergone implantation of percutaneous leads targeting the femoral and/or sciatic nerves and received PNS for up to 60 days.\(^{42}\) Each patient reported 50% or greater reductions in post-amputation pain during the stimulation period (Fig. 2). Two patients were followed for an additional 8 and 16 weeks after lead withdrawal and continued to report 57% and 75% reductions, respectively, in post-amputation pain relative to their pre-PNS baselines.

Collectively, the studies and clinical reports reviewed here suggest that percutaneous PNS can effectively treat multiple aspects of a complex pain state like PAP. Overall, 18/24 (75%) of patients reported substantial (\( \geq 50\% \)) clinically significant relief of both RLP and PLP during the stimulation period, suggesting that the therapy decreased both nociceptive and neuropathic sources of pain. This is consistent with evidence from traditional implanted PNS systems that PNS can attenuate transmission of nociceptive signals from the periphery,\(^{43}\) and attenuate neuropathic pain of various etiologies.\(^{22,24,44}\) In both the nociceptive and neuropathic cases, it is theorized that percutaneous PNS activates spinal segmental inhibitory mechanisms to attenuate pain, such as the gate control mechanism originally proposed by Wall and Melzack,\(^{45}\) to reduce pain during the stimulation period. Of additional interest are the reports that pain relief endured for significant periods of time after the end of the percutaneous PNS stimulation period (pain relief up to one year in 4 of 5, 80%).\(^{39}\) These results suggest a modification of the neuropathic and nocicplastic mechanisms associated with centralized chronic PAP, especially the maladaptive cortical plasticity believed to underlie PLP.\(^{46}\) Percutaneous PNS may enable reversal of aberrant plasticity by modulating painful signals from the periphery, as suggested by review of recent results following nerve blocks for PLP.\(^{46}\) Analysis of studies using cutaneous EMG to provide sensory feedback to the cortex suggests that PNS may, over the course of the therapy period, enable beneficial cortical plasticity by generating non-nociceptive sensory input from the periphery.\(^{47}\) By promoting functional plasticity in response to non-painful sensory input from stimulation, percutaneous PNS therapy may “unwind” nocicplastic mechanisms, thereby altering the chronic pain state. This may manifest as a decrease or absence of pain after withdrawal of the stimulating leads, even in the presence of ongoing nociceptive or neuropathic input.

THE PERCUTANEOUS PNS SYSTEM AND FINE-WIRE COILED LEADS HAVE DEMONSTRATED A STRONG SAFETY PROFILE

The percutaneous PNS system has demonstrated a strong safety profile in numerous studies in chronic and acute pain indications.\(^ {33–37,48–51}\) Consistent with previous studies using percutaneous PNS for shoulder pain, low back pain, and acute post-surgical pain, the most common adverse events reported during the use of percutaneous PNS for post-amputation pain were mild discomfort due to the lead implantation procedure or lead withdrawal, and irritation at the lead exit site or related to adhesive tapes and bandages.\(^ {30,39}\) The rate of suspected lead fracture during treatment of chronic pain following amputation was also consistent with previous reports from a variety of indications (7.5% across 267 leads).\(^ {31–37,49,52,53}\) MR-Conditional lead fragments\(^ {54}\) most commonly occurred during lead withdrawal at or near the tip of the lead, leaving a relatively short length of the 100-micron wire lead. Fragments were observed in situ and no fragment-related sequelae were subsequently reported during follow-up.\(^ {30,39}\) similar to previous reports.\(^ {33,35,37,38}\)

PNS and percutaneous systems were historically associated with risk of infection, but a recent review found that an open-coil lead design significantly reduces this risk. Across 46 studies with over 7,000 coiled and non-coiled leads, the number of infections per lead indwelling time was 25-times lower in coiled leads compared to traditional non-coiled leads used with previous methods of PNS.\(^ {55}\)
PERCUTANEOUS PNS HAS POTENTIAL APPLICATIONS IN POST-SURGICAL AMPUTATION PAIN AND NON-AMPUTEE MILITARY SERVICE MEMBERS AND VETERANS WITH POST-TRAUMATIC PAIN

Multiple studies have evaluated the efficacy and safety of percutaneous PNS in the back and upper and lower extremities for the treatment of acute and chronic pain conditions in addition to post-amputation pain, including for the treatment of chronic neuropathic pain, musculoskeletal pain and postsurgical pain.30–37,48 A recent review of these data found that approximately 89% of patients treated with percutaneous PNS experienced clinically-meaningful (≥ 30%49) pain relief and/or reductions in the interference of pain on daily activities, and 74% experienced substantial (≥ 50%50) benefit, suggesting the therapy has the potential to be employed in patients with a wide range of pain indications.30,35–37,39,48,49,51,53,56–58

Amputation surgeries cause significant post-surgical pain in addition to the potential development of chronic nociceptive, neuropathic, and nociplastic pain. Acute post-amputation pain and post-surgical opioid consumption are key predictors of the development of chronic RLP and PLP,59 and as such post-surgical pain management is an important part of post-amputation care. Pain management strategies have been designed to reduce the risk of chronic post-surgical pain, but current strategies can be ineffective and are often limited by risk of complications (e.g., motor deficit or infection from peripheral nerve catheters) and undesirable side effects (e.g., nausea and sedation from opioid medications).60,61 Recent data suggest that percutaneous PNS can provide pain relief, functional benefit, and reduced opioid consumption in the post-surgical period following total knee replacement and a variety of ambulatory surgeries.31,33,34,38 One ongoing study is evaluating outcomes following amputation (including the incidence of chronic RLP and PLP) in patients treated with percutaneous PNS postsurgically compared to those treated with conventional pain management (Clinicaltrials.gov NCT03484429). Results from an initial case suggest that percutaneous PNS can help manage acute post-amputation pain and may have some impact on post-surgical opioid use.62 Future results from this study will be important to help determine the potential impact of PNS in the acute post-amputation setting.

Amputation is highly visible and debilitating, but traumatic combat injuries can also cause chronic pain without requiring amputation. Nerve injuries are more prevalent than amputation in trauma patients, and a large majority of nerve injuries result in chronic pain.63 Whereas amputations represented just over 3% of the 52,351 reported casualties across all services from 2001–2015,2 up to 24–53% of Service members injured in OIF/OEF had extremity pain.64,65 Because trauma and subsequent amputation result in nerve injuries similar to those in non-amputee nerve trauma patients, data from the successful use of percutaneous PNS in amputees suggest this therapy has the potential to provide significant pain relief and restoration of function following other types of injuries or trauma (both combat- and noncombat-related) that have a high prevalence among military Service members and Veterans.

CONCLUSIONS

Chronic pain in amputees is an important barrier that must be overcome in order to promote functional recovery and effective rehabilitative therapy among injured military Service members and Veterans. Recent evidence suggests that percutaneous PNS may effectively reduce chronic pain following amputation. This therapy fills an unmet need and has the potential to become a standard option to relieve post-amputation pain, and through such pain relief, optimize recovery and restoration of function for Service members and Veterans with amputations. By reducing pain and thereby restoring function in active duty amputees, percutaneous PNS may offer a viable new rehabilitative strategy that could ultimately enable return to duty. Percutaneous PNS may also provide additional benefit for military Service members and Veterans with chronic post-surgical, post-traumatic, or musculoskeletal pain.

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