

PERIPHERAL NERVE STIMULATION

Effective Date: December 1, 2024 Review Dates: 11/22, 5/23, 11/23, 11/24

Date Of Origin: November 23, 2022 Status: Current

Summary of Changes

Additions:

• Scope was broadened to include:

- Transcutaneous Electrical Acustimulation for Hyperemesis (TEA). The corresponding medical policy by that name, No. 91576, is being retired. All contents from that policy have been transcribed to this policy.
- Tonic Motor Activation (TOMAC) peroneal nerve stimulation therapy for restless leg syndrome (RLS) (e.g., Nidra).
- Sparrow Ascent (Spark Biomedical, Inc.) and other devices/stimulators/systems that target nerves in the auricular region and are indicated to reduce symptoms associated with opioid withdrawal was added to the section Percutaneous electrical nerve stimulation (PENS) and percutaneous neuromodulation therapy (PNT)

I. SCOPE

This policy addresses the use of electrical nerve stimulation on peripheral nerves to treat pain, including the following modalities:

- Transcutaneous electrical nerve stimulation (TENS) devices
- Transcutaneous Electrical Acupoint Stimulation (TEAS)
- Percutaneous electrical nerve stimulation (PENS) and percutaneous neuromodulation therapy (PNT) devices
- Permanently implanted peripheral nerve stimulation and peripheral nerve field stimulation devices
- Tonic Motor Activation (TOMAC) peroneal nerve stimulation therapy for restless leg syndrome (RLS)

Related medical policies:

For the use of electrical stimulation to treat other conditions (e.g., incontinence, skin ulcers) or the use of electrical stimulation on non-peripheral nerves (e.g., brain, spinal cord/dorsal column, dorsal root ganglion), see *Priority Health Medical Policy No. 91468 – Stimulation Therapy and Devices*.

For hypoglossal nerve stimulation for the treatment of obstructive sleep apnea, see *Priority Health Medical Policy No. 91333 –Sleep Apnea: Obstructive and Central*

Peripheral Nerve Stimulation

For gastric pacing (gastric pacemaker) and gastric electrical stimulation for treatment of gastroparesis, see *Priority Health Medical Policy No. 91572 – Gastroparesis Testing and Treatment.*

For transcranial magnetic stimulation for treatment of depression, see <u>Priority</u> <u>Health Provider Manual: Transcranial magnetic stimulation (TMS)</u>.

II. POLICY/CRITERIA

A. Transcutaneous electrical nerve stimulators (TENS)

- 1. Use of TENS for any diagnosis for a two-month trial does not require prior authorization.
- 2. Authorization of TENS beyond the two-month initial trial for any diagnosis (except those listed in "3" below) requires documentation of at least two of the following:
 - a. Increased physical activity
 - b. Decreased pain
 - c. Decreased use of analgesics
- 3. Use of TENS for the following low back diagnoses does NOT require prior authorization:
 - Intervertebral disc degeneration
 - Spinal instabilities
 - Sacrococcygeal disorders
 - Dorsopathies
 - Low back pain
 - Dorsalgia
- 4. Transcutaneous electrical nerve stimulators (TENS) include the following (not an all-inclusive list):
 - iRelieve Microcurrent Pain Relief System (Fast Track Technologies, Inc.)
 - StimOnTM Pain Relief System (Gimer Medical Co., Ltd.)
 - TrueRelief (TrueRelief)
 - **BioWaveGo** (Biowave Corporation)

The above devices have been classified by the U.S. Food and Drug Administration (FDA) as <u>Stimulator</u>, <u>Nerve</u>, <u>Transcutaneous</u>, <u>For Pain Relief</u> (Classification Product Code GZJ).

- 5. **Limitations/Exclusions**: The following TENS and TENS-related devices are considered experimental, investigational, or unproven:
 - The Monarch eTNS® [external trigeminal nerve stimulation] System (NeuroSigma Inc.) for treatment of attention-deficit/hyperactivity disorder (ADHD)

B. Transcutaneous Electrical Acupoint Stimulation (TEAS)

- 1. Prescription TEAS devices (e.g., prescription version PrimaBellaTM or ReliefBand devices) are medically necessary for the treatment of hyperemesis gravidarum that is unresponsive to other conservative medical therapy (e.g., change in diet, ginger capsules, vitamin B6).
- 2. Over-the-counter (OTC) disposable TEAS devices, which are used for the treatment of motion sickness, are not considered to be medically necessary.

C. Percutaneous electrical nerve stimulation (PENS) and percutaneous neuromodulation therapy (PNT)

- 1. There is insufficient evidence in the published peer-reviewed literature to support the safety and effectiveness of PENS or PNT as a treatment option for any indication. Therefore, Priority Health considers the use of PENS or PNT NOT medically necessary for ANY indication.
- 2. Percutaneous electrical nerve stimulation (PENS) devices include the following (not an all-inclusive list):
 - Sprint® PNS (SPR Therapeutics, Inc.)
 - Smartpatch PNS (SPR Therapeutics, Inc.)
 - Primary Relief, ANSIStim-PP, First Relief (DyAnsys, Inc.)
 - Deepwave Percutaneous Neuromodulation Pain Therapy System (Biowave Corporation)
 - Vertis PNT (Vertis Neuroscience, Inc.)

The above devices have been classified by the **U.S. Food and Drug Administration (FDA)** as <u>Stimulator, Nerve, Electrical, Percutaneous (Pens), For Pain Relief (Classification Product Code NHI).</u>

• Sparrow Ascent (Spark Biomedical, Inc.) and other devices/stimulators/systems that target nerves in the auricular

region and are indicated to reduce symptoms associated with opioid withdrawal.

The above devices have been classified by the U.S. Food and Drug Administration (FDA) as <u>percutaneous nerve stimulator for opioid</u> withdrawal (Classification Product Code PZR).

D. Permanently implanted peripheral nerve stimulators

1. Restorative neurostimulation

The FDA granted Premarket Approval (PMA) for the **ReActiv8**® **Implantable Neurostimulation System (Mainstay Medical Ltd.)** on June 16, 2020. Priority Health considers this treatment modality/device unproven and not medically necessary due to insufficient evidence of efficacy.

ReActiv8® has been classified by the U.S. Food and Drug Administration (FDA) as <u>Stimulator, neuromuscular, Lower Back Muscles, Totally Implanted for Pain Relief (Classification Product Code QLK).</u>

- 2. A permanently implanted peripheral nerve stimulator may be considered medically necessary only when ALL of the following criteria are met:
 - a. Member has been diagnosed with one or more of the following:
 - Reflex sympathetic dystrophy
 - o Causalgia
 - o Plexus avulsion
 - Operative trauma
 - o Entrapment neuropathies
 - Injection injuries
 - b. There is objective evidence of pathology (e.g., electromyography)
 - c. Member is refractory to one or more of the following conservative therapies:
 - Analgesics
 - o Physical therapy
 - Local injection
 - d. Member exhibits no psychological contraindications
 - e. Member is not addicted to any drug

Peripheral Nerve Stimulation

- f. Member has completed a successful two-week trial of transcutaneous stimulation (resulting in at least a 50% reduction in pain).
- 3. Permanently implanted peripheral nerve stimulators include the following (not an all-inclusive list):
 - Nalu Neurostimulation System (Nalu Medical)
 - StimRouter Neuromodulation System (Bioventus)
 - Neuspera Neurostimulation System (Neuspera Medical Inc.)
 - StimQ Peripheral Nerve Stimulator (Stimwave Technologies, Inc.)

The above devices have been classified by the U.S. Food and Drug Administration (FDA) as <u>Stimulator, Peripheral Nerve, Implanted</u> (Pain Relief) (Classification Product Code GZF).

4. Prior authorization is required.

E. Tonic Motor Activation (TOMAC) peroneal nerve stimulation therapy for restless leg syndrome (RLS)

- 1. There is insufficient evidence in the published peer-reviewed literature to support the safety and effectiveness of TOMAC peroneal nerve stimulation therapy as a treatment option for RLS. Therefore, Priority Health considers the use of TOMAC peroneal nerve stimulation therapy NOT medically necessary for RLS.
- 2. Tonic Motor Activation (TOMAC) peroneal nerve stimulators include the following (may not be an all-inclusive list):
 - NidraTM NTX100 Tonic Motor Activation (TOMAC) System
 (Noctrix Health, Inc.). This Class II device has been classified by the U.S. Food and Drug Administration (FDA) as <u>Stimulator</u>, <u>Nerve</u>, <u>For Restless Legs Syndrome (Classification Product Code QWD)</u>.

III. MEDICAL NECESSITY REVIEW

Required Required

• **Medicare**: Prior authorization is required from the start of the rental period.

Peripheral Nerve Stimulation

- Commercial/Individual, Medicaid: Use of TENS beyond the two-month initial trial for any diagnosis (except those listed in section II. A. 3. above)
- Permanently implanted peripheral nerve stimulators

Not Required

- Commercial/Individual, Medicaid: Use of TENS for any diagnosis for a two-month trial.
- **Commercial/Individual, Medicaid**: Use of TENS for any of the low back diagnoses listed in section II. A. 3. above

IV. APPLICATION TO PRODUCTS

Coverage is subject to member's specific benefits. Group specific policy will supersede this policy when applicable.

- **❖** HMO/EPO: This policy applies to insured HMO/EPO plans.
- **POS:** This policy applies to insured POS plans.
- * PPO: This policy applies to insured PPO plans. Consult individual plan documents as state mandated benefits may apply. If there is a conflict between this policy and a plan document, the provisions of the plan document will govern.
- ASO: For self-funded plans, consult individual plan documents. If there is a conflict between this policy and a self-funded plan document, the provisions of the plan document will govern.
- * INDIVIDUAL: For individual policies, consult the individual insurance policy. If there is a conflict between this medical policy and the individual insurance policy document, the provisions of the individual insurance policy will govern.
- **❖** MEDICARE: Coverage is determined by the Centers for Medicare and Medicaid Services (CMS); if a coverage determination has not been adopted by CMS, this policy applies.
- * MEDICAID/HEALTHY MICHIGAN PLAN: For Medicaid/Healthy Michigan Plan members, this policy will apply. Coverage is based on medical necessity criteria being met and the appropriate code(s) from the coding section of this policy being included on the Michigan Medicaid Fee Schedule located at: http://www.michigan.gov/mdch/0,1607,7-132-2945 42542 42543 42546 42551-159815--,00.html. If there is a discrepancy between this policy and the Michigan Medicaid Provider Manual located at: http://www.michigan.gov/mdch/0,1607,7-132-2945 5100-87572--,00.html, the Michigan Medicaid Provider Manual will govern. If there is a discrepancy or lack of guidance in the Michigan Medicaid Provider Manual, the Priority Health contract with Michigan Medicaid will govern. For Medical Supplies/DME/Prosthetics and Orthotics, please refer to the Michigan Medicaid Fee Schedule to verify coverage.

V. BACKGROUND

Transcutaneous electrical nerve stimulators (TENS)

Transcutaneous electrical nerve stimulation (TENS) is a therapy that uses low voltage electrical current to provide pain relief. A TENS unit consists of a battery-powered device that delivers electrical impulses through **electrodes placed on the surface of the skin**. The electrodes are placed at or near nerves where the pain is located or at trigger points. It may be applied in a variety of settings (in the patient's home, a physician's office, or in an outpatient clinic).

There are two theories about how transcutaneous electrical nerve stimulation (TENS) works. One theory is that the electric current stimulates nerve cells that block the transmission of pain signals, modifying your perception of pain. The other theory is that nerve stimulation raises the level of endorphins, which are the body's natural pain-killing chemical. The endorphins then block the perception of pain.

The Monarch eTNS [external trigeminal nerve stimulation] System is designed to provide a nonpharmaceutical treatment option for children with attention-deficit/hyperactivity disorder (ADHD) during sleep without the need for device implantation. The U.S. Food and Drug Administration (FDA) granted a de novo (DEN) classification (DEN180041) for the Monarch eTNS System (NeuroSigma Inc.) under product code QGL (transcutaneous nerve stimulator for ADHD). The Monarch eTNS System is a class II device, regulated under Code of Federal Regulations (CFR) 21 CFR 882.5898. Treatment takes place in the patient's home environment. The Monarch eTNS System is a prescription-only, noninvasive, therapeutic device intended for children ages 7 to 12 years diagnosed with ADHD who are not taking medications. Before the patient goes to sleep at night, a new electrical patch is adhered to clean unbroken skin in the midline of the patient's forehead directly above the eyebrows. The patch is connected to the generator by a conductive wire. The pulse generator, operating on preset parameters—except for amplitude, which can be adjusted in 0.2 milliamp (mA) increments and then locked by the caretaker under direction of the prescribing physician—delivers low-level current via the patch to the patient's right and left supraorbital and supratrochlear branches of the trigeminal nerve. Prescribed duration of use while sleeping ranges from 7 to 9 hours. According to the American Academy of Pediatrics (AAP) Clinical Practice Guideline for the Diagnosis, Evaluation, and Treatment of Attention-Deficit/Hyperactivity Disorder in Children and Adolescents: "To date, there is no long-term safety and efficacy evidence for eTNS [external trigeminal nerve stimulation]. Overall, the current evidence supporting [tx] of ADHD with eTNS is sparse and in no way approaches the robust strength of evidence documented for established medication and behavioral [tx] for ADHD; therefore, it cannot be recommended as a [tx] of

ADHD without considerably more extensive study on its efficacy and safety" (p. 13).

Transcutaneous Electrical Acupoint Stimulation (TEAS) for hyperemesis gravidarum

Up to 90% of pregnant women experience nausea and vomiting. When prolonged or severe, this is known as hyperemesis gravidarum (HG), which can, in individual cases, be life threatening. The etiology of HG is unknown in most cases, although some biological, physiological and psychological as well as sociocultural factors are thought to be contributory factors. Risk factors for HG include multiple pregnancy, nulliparity, obesity, metabolic disturbances, a history of HG in a previous pregnancy, trophoblastic disorders, psychological disorders (for example, eating disorders such as anorexia nervosa or bulimia) and a history of migration. For initial management, dietary and lifestyle advice is often sufficient to ameliorate symptoms and improve quality of life. TEAS devices emit a low-level electrical current across two small electrodes on their underside, stimulating the median nerve (an acupuncture point).

Percutaneous Electrical Nerve Stimulation (PENS), Percutaneous Neuromodulation Therapy (PNT) and Percutaneous Electrical Nerve Field Stimulation (PENFS)

Percutaneous electrical nerve stimulation (PENS) and percutaneous neuromodulation therapy (PNT) are therapies that combine the features of electroacupuncture and transcutaneous electrical nerve stimulation (TENS).

Percutaneous electrical nerve stimulation (PENS) involves the use of thin filiform needle electrodes that are placed percutaneously near a peripheral nerve. It may also involve the use of a needle-like introducer that inserts an electrode near a peripheral nerve. An electrical current drawn from an external pulse generator is delivered to the area, aiming to interfere with pain sensation. PENS devices are temporary and do not require invasive procedures to administer.

Percutaneous electrical nerve field stimulation (PENFS) differs from PENS in that with PENFS, a "field" of pain is targeted, instead of targeting a specific nerve.

Percutaneous neuromodulation therapy (PNT) is a variant of PENS in which fine filament electrode arrays are placed near the area that is causing pain. Some use the terms PENS and PNT interchangeably. It is proposed that PNT inhibits pain transmission by creating an electrical field that hyperpolarizes C-fibers, thus preventing action potential propagation along the pain pathway.

Permanently implanted peripheral nerve stimulators

Restorative neurostimulation

The FDA granted Premarket Approval (PMA) for the **ReActiv8®** Implantable **Neurostimulation System (Mainstay Medical Ltd.)**. This device is indicated for bilateral stimulation of the L2 medial branch of the dorsal ramus as it crosses the transverse process at L3 as an aid in the management of intractable chronic low back pain associated with multifidus muscle dysfunction, as evidenced by imaging or physiological testing in adults who have failed therapy including pain medications and physical therapy and are not candidates for spine surgery.

ReActiv8® was formally evaluated by Priority Health's **Technology Assessment Committee (TAC)** on August 30, 2023. It is the company's position that there is insufficient evidence in the published peer reviewed scientific literature to support the efficacy of restorative neurostimulation for the treatment of chronic low back pain. Additional larger studies comparing restorative neurostimulation to standard of care and current alternative treatments are needed to demonstrate safety and efficacy for this modality. Therefore, Priority Health considers this device/treatment modality unproven and not medically necessary due to insufficient evidence of efficacy.

Peripheral nerve stimulation

Peripheral nerve stimulation (PNS) involves surgical insertion of an electrode along a specific peripheral nerve determined to be responsible for regional pain. The electrode is connected to a lead that is tunneled to a receiver unit located within a subcutaneous pocket. Electrical impulses generated by a stimulator attached to the skin overlying the receiver are transmitted along the electrode to the peripheral nerve, thereby blocking or masking pain sensation. A therapeutic trial may be attempted by placement of a temporary electrode to determine if nerve stimulation leads to significant therapeutic analgesia - by at least 50 %. Individuals that experience significant pain relief may then be eligible for permanent implantation.

Peripheral Nerve Field Stimulation (PNFS)

Subcutaneous stimulation (peripheral nerve field stimulation/PNFS) is a novel neuromodulation modality that has increased in its utilization during the past decade. It consists of introducing a lead in the subdermal level to stimulate the small nerve fibers in that layer. Unlike other neuromodulation techniques including direct peripheral nerve stimulation, spinal cord stimulation (SCS), or deep brain stimulation, the precise target is not identified.

VI. CODING INFORMATION

TENS

Transcutaneous Electrical Stimulator (TENS)

ICD-10 Codes that <u>may</u> apply:

• No prior auth required for this indication

No prior auth for first 2 months trial for any indication for commercial and Medicaid.

Prior auth required for <u>Medicare</u> for all indications from 1st months rental		
• •		
B02.0	Zoster encephalitis	
B02.23	Postherpetic polyneuropathy	
B02.29	Other postherpetic nervous system involvement	
E08.40 – E08.42	Diabetes mellitus due to underlying condition with neurological complications	
E09.40 – E09.42	Drug or chemical induced diabetes mellitus with neurological complications	
E10.40 - E10.49	Type 1 diabetes mellitus with neurological complications	
E10.610	Type 1 diabetes mellitus with diabetic neuropathic arthropathy	
E10.65	Type 1 diabetes mellitus with hyperglycemia	
E11.40 - E11.49	Type 2 diabetes mellitus with neurological complication	
E11.610	Type 2 diabetes mellitus with diabetic neuropathic arthropathy	
E11.65	Type 2 diabetes mellitus with hyperglycemia	
E13.40	Other specified diabetes mellitus with diabetic neuropathy, unspecified	
E13.41 - E13.49	Other specified diabetes mellitus with neurological complication	
G54.8	Other nerve root and plexus disorders	
G55	Nerve root and plexus compressions in diseases classified	
	elsewhere	
G57.70 - G57.72	Causalgia of lower limb	
G57.80 - G57.82	Other specified mononeuropathies of left lower limb	
G57.90 - G57.92	Unspecified mononeuropathy of lower limb	
G58.8	Other specified mononeuropathies	
G58.9	Mononeuropathy, unspecified	
G59	Mononeuropathy in diseases classified elsewhere	
G89.0	Central pain syndrome	
G89.21 - G89.29	Chronic pain	
G89.4	Chronic pain syndrome	
G90.50 - G90.59	Complex regional pain syndrome I	
G99.0	Autonomic neuropathy in diseases classified elsewhere	
M43.20 - M43.28	Fusion of spine	
M43.8x9	Other specified deforming dorsopathies, site unspecified	
M51.36*	Other intervertebral disc degeneration, lumbar region	
M51.37◆	Other intervertebral disc degeneration, lumbosacral region	
M53.2x7◆	Spinal instabilities, lumbosacral region	
M53.2x8◆	Spinal instabilities, sacral and sacrococcygeal region	
M53.3◆	Sacrococcygeal disorders, not elsewhere classified	
M53.80	Other specified dorsopathies, site unspecified	
2	1	



M53.84	Other specified dorsopathies, thoracic region
M53.85	Other specified dorsopathies, thoracolumbar region
M53.86◆	Other specified dorsopathies, lumbar region
M53.87◆	Other specified dorsopathies, lumbosacral region
M53.88◆	Other specified dorsopathies, sacral and sacrococcygeal region
M53.9	Dorsopathy, unspecified
M54.5◆	Low back pain
M54.89	Other dorsalgia
M54.9◆	Dorsalgia, unspecified
СРТ/НС	PCS Codes:
97014	Application of a modality to one or more areas; electrical stimulation
<i>)</i> / 01 1	(unattended) (No Auth) (Not covered for Medicare)
97032	Application of a modality to one or more areas; electrical stimulation (manual),
71032	each 15 minutes (No Auth)
	Cach 13 minutes (110 1100)
G0283	Electrical stimulation (unattended), to one or more areas for indication(s) other
	than wound care, as part of a therapy plan of care (Medicare only) (No Auth)
A4558	Conductive gel or paste, for use with electrical device (e.g., TENS, NMES),
	per oz (No Auth)
A4595	Electrical stimulator supplies, 2 lead, per month, (e.g. TENS, NMES)
	(No Auth)
A4630	Replacement batteries, medically necessary, transcutaneous electrical
	stimulator, owned by patient (No Auth) (Not covered by Priority Medicaid)
E0720	Transcutaneous electrical nerve stimulation (TENS) device, two lead, localized
	stimulation
E0730	Transcutaneous electrical nerve stimulation (TENS) device, four or more leads,
	for multiple nerve stimulation
E0731	Form-fitting conductive garment for delivery of TENS or NMES (with
	conductive fibers separated from the patient's skin by layers of fabric)
	(Covered for Medicare, Medicaid ONLY)
Not cover	
0278T	Transcutaneous electrical modulation pain reprocessing (eg, scrambler
	therapy), each treatment session (includes placement of electrodes)
A4541	Monthly supplies for use of device coded at E0733
A4542	Supplies and accessories for external upper limb tremor stimulator of the
	peripheral nerves of the wrist
A4543	Supplies for transcutaneous electrical nerve stimulator, for nerves in the
	auricular region, per month
A4544	Electrode for external lower extremity nerve stimulator for restless legs
	syndrome
A4545	Supplies and accessories for external tibial nerve stimulator (e.g., socks, gel
	pads, electrodes, etc.), needed for one month
E0733	Transcutaneous electrical nerve stimulator for electrical stimulation of the
	trigeminal nerve



Peripheral Nerve Stimulation

E0734 E0721	External upper limb tremor stimulator of the peripheral nerves of the wrist Transcutaneous electrical nerve stimulatory, stimulates nerves in the auricular region
E0737	Transcutaneous tibial nerve stimulator, controlled by phone application
E0743	External lower extremity nerve stimulator for restless legs syndrome, each

Transcutaneous Electrical Acupoint Stimulation (TEAS) for hyperemesis gravidarum

ICD-10 Codes that may support medical necessity:

- O21.0 Mild hyperemesis gravidarum
- O21.1 Hyperemesis gravidarum with metabolic disturbance
- O21.2 Late vomiting of pregnancy
- O21.8 Other vomiting complicating pregnancy
- O21.9 Vomiting of pregnancy, unspecified

CPT/HCPCS Codes:

E0765 FDA approved nerve stimulator, with replaceable batteries, for treatment of nausea and vomiting (Not covered for Priority Health Medicaid/Healthy Michigan Plan members)

PENS

Percutaneous Electrical Nerve Stimulation (PENS) and Percutaneous Neuromodulation Therapy (PNT) Not Covered:

97813	Acupuncture, 1 or more needles; with electrical stimulation, initial 15 minutes
	of personal one-on-one contact with the patient
97814	Acupuncture, 1 or more needles; with electrical stimulation, each additional 15
	minutes of personal one-on-one contact with the patient, with re-insertion of
	needle(s) (List separately in addition to code for primary procedure)
64999	Unlisted procedure, nervous system (Explanatory notes must accompany
	claims billed with unlisted codes.)
0720T	Percutaneous electrical nerve field stimulation, cranial nerves, without

0720T Percutaneous electrical nerve field stimulation, cranial nerves, without implantation

C0807 Norwa stimulator, percutaneous, peripheral (e.g., sprint peripheral nerv

C9807 Nerve stimulator, percutaneous, peripheral (e.g., sprint peripheral nerve stimulation system), including electrode and all disposable system components, non-opioid medical device (must be a qualifying medicare non-opioid medical device for post-surgical pain relief in accordance with section 4135 of the caa, 2023) (Covered for Medicare and Medicaid)

Peripherally Implanted Nerve Stimulator



Peripheral Nerve Stimulation

ICD-10 Codes that may apply:		
G54.8	Other nerve root and plexus disorders	
G54.9	Nerve root and plexus disorders Nerve root and plexus disorder, unspecified	
G55	Nerve root and plexus compressions in diseases classified	
033	elsewhere	
G56.40 - G56.42	Causalgia of upper limb	
G56.80 - G56.82	Other specified mononeuropathies	
G57.70 - G57.72	Causalgia of lower limb	
G57.80 - G57.82	Other specified mononeuropathies	
G58.0	Intercostal neuropathy	
G58.7	Mononeuritis multiplex	
G58.8	Other specified mononeuropathies	
G89.0	Central pain syndrome	
G89.21	Chronic pain due to trauma	
G89.22	Chronic post-thoracotomy pain	
G89.28	Other chronic postprocedural pain	
G89.29	Other chronic pain	
G89.4	Chronic pain syndrome	
G90.50 - G90.59	Complex regional pain syndrome I	
M53.80	Other specified dorsopathies, site unspecified	
M53.84	Other specified dorsopathies, thoracic region	
M53.85	Other specified dorsopathies, thoracolumbar region	
M53.9	Dorsopathy, unspecified	
M54.5	Low back pain	
M54.89	Other dorsalgia	
M54.9	Dorsalgia, unspecified	
W134.9	Dorsaigia, unspectfied	
CPT/HCPCS Codes:		
	s implantation of neurostimulator electrodes; peripheral nerve	
(excludes sa	* * *	
`	,	
64575 Incision for	implantation of neurostimulator electrodes; peripheral nerve	
(excludes sa	cral nerve)	
64590 Insertion or	replacement of peripheral, sacral, or gastric neurostimulator pulse	
generator or	receiver, requiring pocket creation and connection between	
electrode arr	ray and pulse generator or receiver	
	removal of peripheral, sacral, or gastric neurostimulator pulse	
generator or	receiver, with detachable connection to electrode array (No Auth)	
	replacement of percutaneous electrode array, peripheral nerve, with	
	eurostimulator, including imaging guidance, when performed;	
imitial alaatu		

Insertion or replacement of percutaneous electrode array, peripheral nerve, with integrated neurostimulator, including imaging guidance, when performed; each additional electrode array (List separately in addition to code for primary

Revision or removal of neurostimulator electrode array, peripheral nerve, with

initial electrode array

integrated neurostimulator

procedure)

64597

64598

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95970	Electronic analysis of implanted neurostimulator pulse generator/transmitter
	(e.g., contact group[s], interleaving, amplitude, pulse width, frequency [Hz], on/off cycling, burst, magnet mode, dose lockout, patient selectable
	parameters, responsive neurostimulation, detection algorithms, closed loop
	parameters, and passive parameters) by physician or other qualified health care
	professional; with brain, cranial nerve, spinal cord, peripheral nerve, or sacral
	nerve, neurostimulator pulse generator/transmitter, without programming (No
	Auth)
95971	Electronic analysis of implanted neurostimulator pulse generator/transmitter
	(e.g., contact group[s], interleaving, amplitude, pulse width, frequency [Hz], on/off cycling, burst, magnet mode, dose lockout, patient selectable
	parameters, responsive neurostimulation, detection algorithms, closed loop
	parameters, and passive parameters) by physician or other qualified health care
	professional; with simple spinal cord or peripheral nerve (eg, sacral nerve)
	neurostimulator pulse generator/transmitter programming by physician or other
0.5072	qualified health care professional (No Auth)
95972	Electronic analysis of implanted neurostimulator pulse generator/transmitter (eg, contact group[s], interleaving, amplitude, pulse width, frequency [Hz],
	on/off cycling, burst, magnet mode, dose lockout, patient selectable
	parameters, responsive neurostimulation, detection algorithms, closed loop
	parameters, and passive parameters) by physician or other qualified health care
	professional; with complex spinal cord or peripheral nerve (eg, sacral nerve)
	neurostimulator pulse generator/transmitter programming by physician or other
	qualified health care professional. (No Auth)
A4438	Adhesive clip applied to the skin to secure external electrical nerve stimulator
~	controller, each
C1767	Generator, neurostimulator (implantable), nonrechargeable
C1778	Lead, neurostimulator (implantable)
C1787 C1816	Patient programmer, neurostimulator Pageiver and/or transmitter, neurostimulator (implentable)
C1810	Receiver and/or transmitter, neurostimulator (implantable) Generator, neurostimulator (implantable), with rechargeable battery and
C1620	charging system
C1822	Generator, neurostimulator (implantable), high frequency, with rechargeable
	battery and charging system
C1883	Adapter/ extension, pacing lead or neurostimulator lead
C1897	Lead, neurostimulator test kit (implantable)
L8679	Implantable neurostimulator, pulse generator, any type
L8680	Implantable neurostimulator electrode, each
L8681	Patient programmer (external) for use with implantable programmable
	neurostimulator pulse generator
L8689	External recharging system for battery (internal) for use with implantable
T 0 60 =	neurostimulator
L8695	External recharging system for battery (external) for use with implantable
	neurostimulator, replacement only
	(L codes not separately paid under APC payment arrangements)

VII. REFERENCES

- 1. Hayes, Inc. Evolving Evidence Review. ReActiv8 Implantable
 Neurostimulation System (Mainstay Medical Ltd.) for Chronic Low Back
 Pain. Hayes, Inc.; May 20, 2022.
- 2. Hayes, Inc. Health Technology Assessment. **Transcutaneous Electrical Nerve Stimulation for Chronic Low Back Pain**. Hayes, Inc.; September 21, 2018.
- 3. Hayes, Inc. Health Technology Assessment. **Percutaneous Electrical Nerve Stimulation for Treatment of Low Back Pain**. Hayes, Inc.; January 10, 2019.
- 4. Hayes, Inc. Health Technology Assessment. **Peripheral Nerve Field Stimulation for Treatment of Chronic Low Back Pain**. Hayes, Inc.; April 22, 2021.
- 5. Hayes, Inc. Health Technology Assessment. **Percutaneous Peripheral Nerve Stimulation for Treatment of Chronic Pain**. Hayes, Inc.; May 5, 2022.
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