

2017 Ambulatory Surgery Center Medicare Reimbursement Guide for Spinal Cord Stimulation

For support, please contact: reimbursement@nuvectramed.com | 1-844-727-7897

Device	Model	CPT ¹ L Code ²	Description	Multiple Procedure Discounting	Payment Indicator ³	Payment⁴
Percutaneous	s Leads a	nd Extensio	ns			
8-electrode lead	1081 1084 1086	63650	Percutaneous implantation of neurostimulator electrode array, epidural.	No	J8	\$4,422
12-electrode lead	1121 1124 1126	63661	Removal of spinal neurostimulator electrode percutaneous array(s), including fluoroscopy, when performed. (Do not report 63661 when removing or replacing a temporary percutaneously placed array for an external generator.)	No	G2	\$788
Extension	5208 5212	63663	Revision including replacement, when performed, of spinal neurostimulator electrode percutaneous array(s), including fluoroscopy, when performed. (Do not report 63663 when removing or replacing a temporary percutaneously placed array for an external generator.)(Do not report 63663 in conjunction with 63661, 63662 for the same spinal level.)	No	J8	\$4,488
Paddle Leads						
Multi-Midline Paddle	3000	63655	Laminectomy for implantation of neurostimulator electrodes, plate/paddle, epidural.	No	J8	\$14,675
2x6 Paddle	3101	63662	Removal of spinal neurostimulator electrode plate/paddle(s) placed via laminotomy or laminectomy, including fluoroscopy, when performed.	No	G2	\$1,453
		63664	Revision including replacement, when performed, of spinal neurostimulator electrode plate/paddle(s) placed via laminotomy or laminectomy, including fluoroscopy, when performed. (Do not report 63664 in conjunction with 63661, 63662 for the same spinal level.)	No	J8	\$13,669
Stimulators						
Stimulator (3x8)	2408	63685 L8679	Insertion or replacement of spinal neurostimulator pulse generator or receiver, direct or inductive coupling. (Do not report 63685 in conjunction with 63688 for the same pulse generator or receiver.) Implantable neurostimulator pulse generator, any type.	No	J8	\$23,148
Stimulator (2x12)	2412	63688	Revision or removal of implanted spinal neurostimulator pulse generator or receiver.	No	A2	\$1,453
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Multiple



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Programmers and Chargers									
Pocket Programmer	4100	L8681	Patient programmer (external) for use with implantable programmable neurostimulator pulse generator, replacement only. Check with local carrier on coverage for replacement.	N/A	N/A	Check with local carrier			
Patient Programmmer Charger	4200	L8689	External recharging system for battery (internal) for use with implantable neurostimulator, replacement only. Check with local carrier on coverage for replacement.	N/A	N/A	Check with local carrier			
Clinician Programmer	4500	95972*	Electronic analysis of implanted neurostimulator pulse generator system (eg, rate, pulse amplitude, pulse duration, configuration of wave form, battery status, electrode selectability, output modulation, cycling, impedance and patient compliance measures); complex spinal cord, or peripheral (ie, peripheral nerve, sacral nerve, neuromuscular) neurostimulator pulse generator/transmitter, with intraoperative or subsequent programming. *This service should only be billed if it is performed by a physician or physician-supervised personnel. A physician should not bill if the service is performed entirely by, or under the direction of, a manufacturer representative without payor consent. Contact local payor with any questions.	N/A	N/A	Check with local carrier			

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- (2) HCPCS II codes (L-codes) may be used by ambulatory surgery centers for billing to non-Medicare payers. Check with local carrier.
- (3) J8: Device-intensive procedure; paid at adjusted rate.
 - G2: Non office-based surgical procedure added in CY 2008 or later; payment based on OPPS relative payment weight.
 - A2: Surgical procedure on ASC list in CY 2007; payment based on OPPS relative payment weight.
- (4) Medicare 2017 base rates without geographical adjustments. Actual payment will vary based on the maximum allowances less any applicable deductibles, co-insurance, etc.

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This document is intended to provide reimbursement assistance only where products have been used according to their FDA-approved or cleared indications. Where reimbursement is being requested in conjunction with use of a product that is inconsistent with, or not expressly granted in, the FDA-approved labeling (which may be found in the clinician's manual, user's guide or directions for use), please consult your billing personnel or the payor for instructions on the proper handling of this type of claim. Some payors may restrict such claims or services. Contact your MAC or other payor for any questions regarding coverage, coding and payment.

Brief Summary: Product Technical Manuals and Information for Prescribers (IFP) must be consulted prior to use of this product.

Indications for Use: The Algovita® Spinal Cord Stimulation (SCS) System is indicated as an aid in the management of chronic intractable pain of the trunk and/or limbs, including unilateral or bilateral pain. Contraindications: Diathermy, patients who are poor surgical candidates. Warnings/Precautions: Strong electromagnetic interference (eg, electrocautery, RF or microwave ablation, or MRI) can result in serious patient injury or death, unexpected stimulation, or device malfunction or damage. Rupture or piercing of the neurostimulator many result in severe burns. Safety and effectiveness of SCS have not been established for pediatric patients, for use during pregnancy, or for use with nursing patients. Adverse Events: may include painful stimulation or loss of pain relief, hardware malfunction or migration, allergic response and surgical risks, such as infection, or additional surgery. For full prescribing information, please call Nuvectra at 1.844.727.7897 and/or consult Nuvectra's website at www.nuvectramed.com. Rx Only. February 2017.

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