

Clinical Policy: Diaphragmatic/Phrenic Nerve Stimulation

Reference Number: LA.CP.MP.203

Date of Last Revision: 08/2425

Revision Log

See <u>Important Reminder</u> at the end of this policy for important regulatory and legal information.

Description

Diaphragmatic/phrenic nerve stimulation, also referred to as diaphragm pacing, is a treatment option used to eliminate or reduce the need for ventilator support in those with chronic ventilatory insufficiency or failure due to bilateral paralysis or severe paresis of the diaphragm. Diaphragmatic/phrenic nerve stimulation uses the phrenic nerves to signal the diaphragm muscles to contract rhythmically and produce breathing through electrical stimulation. §

Policy/Criteria

- I. It is the policy of Louisiana Healthcare Connections that diaphragmatic/phrenic nerve stimulation with the Mark IV[™] Breathing Pacemaker System or the Spirit Diaphragm Pacing Transmitter is **medically necessary** when all of the following are met:
 - A. Stimulation is used as an alternative to mechanical ventilation for an individual with severe, chronic respiratory failure due to one of the following:
 - 1. Upper cervical spinal cord injury (at or above the C3 vertebral level);
 - 2. Central alveolar hypoventilation disorder;
 - B. Diaphragm movement with stimulation is visible under fluoroscopy;
 - C. Intact and sufficient function in the phrenic nerve, lungs, and diaphragm;
 - D. Stimulation of the diaphragm either directly or through the phrenic nerve results in sufficient muscle activity to accommodate independent breathing without the support of a ventilator;
 - E. Normal chest anatomy, a normal level of consciousness, and the ability to participate in and complete the training and rehabilitation associated with the use of the device.
- **II.** It is the policy of Louisiana Healthcare Connections that diaphragmatic/phrenic nerve stimulation with the NeuRx RA/4 Diaphragm Pacing System® is **medically necessary** when provided in accordance with the Humanitarian Device Exemption (HDE) specifications of the U.S Food and Drug Administration when all of the following are met:
 - A. Stimulation is used as an alternative to mechanical ventilation for an individual with severe, chronic respiratory failure due to one of the following:
 - 1. Amyotrophic lateral sclerosis (ALS);
 - a. Age 21 years or older;
 - b. Experiencing chronic hypoventilation but not progressed to forced vital capacity (FVC) less than 45% predicted;
 - c. Diaphragm movement with stimulation is visible under fluoroscopy or by other radiographic techniques such as ultrasound;
 - d. Intact and sufficient function in the phrenic nerve, lungs, and diaphragm.
 - 2. Upper cervical spinal cord injury (at or above the C3 vertebral level);
 - a. Age 18 years or older;
 - b. Diaphragm movement with stimulation is visible under fluoroscopy or by other radiographic techniques such as ultrasound;



- c. Stimulation of the diaphragm will allow the individual to breathe without the assistance of a mechanical ventilator for at least four continuous hours a day;
- d. Intact and sufficient function in the phrenic nerve, lungs, and diaphragm.
- **III.** It is the policy of Louisiana Healthcare Connections that there is insufficient evidence to support the safety and efficacy of diaphragmatic/phrenic nerve stimulation for any other conditions, including but not limited to, central sleep apnea.

Background

Diaphragmatic/phrenic nerve stimulator devices are indicated for certain ventilator-dependent individuals who lack voluntary control of their diaphragm muscles to enable independent breathing without the assistance of a mechanical ventilator. §

Avery Diaphragm Pacing System (Avery Biomedical Device, Inc.)

The Avery Diaphragm Pacing System includes receivers and electrodes that are surgically implanted. An electrode is placed under the phrenic nerve and is connected to a radiofrequency receiver, which is implanted under the skin.³

The different types of Avery systems include the Mark IV Breathing Pacemaker System and the Spirit Diaphragm Pacing System.³ The Mark IV Breathing Pacemaker System is a diaphragmatic/phrenic stimulator system approved for use by the FDA in the United States. The device is approved "for persons who require chronic ventilatory support because of upper motor neuron respiratory muscle paralysis (RMP) or because of central alveolar hypoventilation (CAH) and whose remaining phrenic nerve, lung, and diaphragm function is sufficient to accommodate electrical stimulation." In 2019, the Spirit Diaphragm Pacing Transmitter received full FDA approval for the use of this system for patients who have functional lungs and diaphragm muscle and who have an intact phrenic nerve. 3.5.6

NeuRx RA/4 Diaphragm Pacing System® (Synapse Biomedical, Inc.)

The NeuRx RA/4 Diaphragm Pacing System® includes implanted intramuscular diaphragm electrodes, which are connected to an external stimulator.²

The United States Food and Drug Administration (FDA) approval for distribution of the NeuRx DPS® (Synapse Biomedical, Inc., Oberlin, OH) was granted under a Humanitarian Device Exemption (HDE) on June 17, 2008. The FDA-approved indications are: "For use in patients with stable, high spinal cord injuries with stimulatable diaphragms, but lack control of their diaphragms." The device is indicated to allow the patients to breathe without the assistance of a mechanical ventilator for at least four continuous hours a day and is for use only in patients 18 years of age or older. This FDA approval is subject to the manufacturer developing an acceptable method of tracking device implantation to individual patient recipients.¹

In 2011, the FDA approved the NeuRx RA/4 Diaphragm Pacing System® as a humanitarian-use device (HUD) in amyotrophic lateral sclerosis (ALS) following the submission of a humanitarian device exemption (HDE) application. The FDA approved indications are: "For use in amyotrophic lateral sclerosis (ALS) patients with a stimulatable diaphragm (both right and left portions) as demonstrated by voluntary contraction or phrenic nerve conduction studies, and who



are experiencing chronic hypoventilation (CH), but not progressed to an FVC less than 45% predicted. For use only in patients 21 years of age or older."^{2(p.1)}

In 2023, NeuRx RA/4 Diaphragm Pacing System received full FDA approval for use in "patients with stable, high spinal cord injuries with stimulatable diaphragms, but who lack control of their diaphragms." ¹⁴

Remedé System (ZOLL® Medical Corporation)

Although the Remedé System was approved as a treatment option for moderate to severe central sleep apnea in adults by the FDA in 2017, there is insufficient evidence to determine the safety and effectiveness of this system. ^{9,15} Additional high-quality studies are recommended to evaluate the clinical significance and long-term safety and efficacy of the Remedé System for central sleep apnea. ⁹

Avery Diaphragm Pacing System (Avery Biomedical Device, Inc.)

The Avery Diaphragm Pacing System includes receivers and electrodes that are surgically implanted and includes an external transmitter worn over the implanted receivers.³ The different types of Avery systems include the Mark IV Breathing Pacemaker System and the Spirit Diaphragm Pacing System.³ The Mark IV Breathing Pacemaker System is a diaphragmatic/phrenic stimulator system approved for use by the FDA in the United States. The device is approved "for persons who require chronic ventilatory support because of upper motor neuron respiratory muscle paralysis (RMP) or because of central alveolar hypoventilation (CAH) and whose remaining phrenic nerve, lung, and diaphragm function is sufficient to accommodate electrical stimulation." In 2019, the Spirit Diaphragm Pacing Transmitter received full FDA approval for the use of this system for patients who have functional lungs and diaphragm muscle and who have an intact phrenic nerve. ^{3,5,6}

Coding Implications

This clinical policy references Current Procedural Terminology (CPT®). CPT® is a registered trademark of the American Medical Association. All CPT codes and descriptions are copyrighted 20232024, American Medical Association. All rights reserved. CPT codes and CPT descriptions are from the current manuals and those included herein are not intended to be all-inclusive and are included for informational purposes only. Codes referenced in this clinical policy are for informational purposes only and may not support medical necessity. Inclusion or exclusion of any codes does not guarantee coverage. Providers should reference the most up-to-date sources of professional coding guidance prior to the submission of claims for reimbursement of covered services.

NOTE: Coverage is subject to each requested code's inclusion on the corresponding LDH fee schedule. Non-covered codes are denoted (*) and are reviewed for Medical Necessity for members under 21 years of age on a per case basis.

CPT®*	Description
Codes	
64575	Open implantation of neurostimulator electrode array; peripheral nerve (excludes sacral nerve)
64580*	Open implantation of neurostimulator electrode array; neuromuscular



CPT®*	Description
Codes	
64590	Insertion or replacement of peripheral, sacral, or gastric neurostimulator pulse
	generator or receiver, requiring pocket creation and connection between electrode
	array and pulse generator or receiver
64595	Revision or removal of peripheral, sacral, or gastric neurostimulator pulse generator
	or receiver, with detachable connection to electrode array

HCPCS ®* Codes	Description			
C1778	Lead, neurostimulator (implantable)			
C1816*	Receiver and/or transmitter, neurostimulator (implantable)			
L8680*	Implantable neurostimulator electrode, each			
L8681*	Patient programmer (external) for use with implantable programmable			
	neurostimulator pulse generator, replacement only			
L8682*	Implantable neurostimulator radiofrequency receiver			
L8683*	Radiofrequency transmitter (external) for use with implantable neurostimulator			
	radiofrequency receiver			
<u>L8685*</u>	Implantable neurostimulator pulse generator, single array, rechargeable, includes			
	extension			
<u>L8686*</u>	Implantable neurostimulator pulse generator, single array, nonrechargeable,			
	<u>includes extension</u>			
<u>L8687*</u>	Implantable neurostimulator pulse generator, dual array, rechargeable, includes extension			
1.00004				
L8688*	Implantable neurostimulator pulse generator, dual array, nonrechargeable, includes extension			
1.0000*				
L8689*	External recharging system for battery (internal) for use with implantable neurostimulator, replacement only			
L8695	External recharging system for battery (external) for use with implantable			
	neurostimulator, replacement only			
L8696	Antenna (external) for use with implantable diaphragmatic/phrenic nerve			
	stimulation device, replacement, each			

Reviews, Revisions, and Approvals	Revision Date	Approval Date	Effective Date
Converted Corporate policy to local policy	2/22	4/14/22	
Annual review. Criteria II.A.1.c. and Criteria II.A.2.b. updated	01/23	4/10/23	
to include "or by other radiographic techniques such as			
ultrasound" in addition to fluoroscopy. Background updated to			
include U.S. Food and Drug Administration premarket approval			
information regarding the Avery Spirit Diaphragm Pacing			
Transmitter. ICD-10 codes removed. Removed codes not on			
LDH fee schedule: CPT 64575, 64580, 64595 and HCPCS			
C1816, L8680, L8681, L8682, L8683, L8689. References			
reviewed and updated.			



Reviews, Revisions, and Approvals	Revision Date	Approval Date	Effective Date
Annual review. Product name updates in criteria II. and in	09/23	11/27/23	
background with no clinical significance. References reviewed			
and updated. Note for non-covered codes added.			
Annual review. Criteria I. updated to include the Spirit	8/24	10/23/24	11/22/24
Diaphragm Pacing Transmitter. Background updated to include			
information regarding full FDA approval of the Spirit			
Diaphragm Pacing Transmitter. References reviewed and			
updated. Reviewed by external specialist.			
Annual review. Description updated with no clinical	08/25		
significance. Background updated to include information			
regarding full FDA approval of NeuRx RA/4 Diaphragm			
Pacing System and added section regarding the Remedé			
System. Added codes L8685, L8686, L8687, L8688 to HCPCs			
Codes table. Coding reviewed. References reviewed and			
<u>updated.</u>			

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Important Reminder

This clinical policy has been developed by appropriately experienced and licensed health care professionals based on a review and consideration of currently available generally accepted standards of medical practice; peer-reviewed medical literature; government agency/program approval status; evidence-based guidelines and positions of leading national health professional organizations; views of physicians practicing in relevant clinical areas affected by this clinical policy; and other available clinical information. LHCC makes no representations and accepts no liability with respect to the content of any external information used or relied upon in developing this clinical policy. This clinical policy is consistent with standards of medical practice current at the time that this clinical policy was approved.

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