

# PRESCRIPTION AND WRITTEN ORDER

(High Frequency Chest Wall Oscillation E0483)

FAX: 866.569.1912

First Name:		Last Name:				
Address:						
City:		State: Zip:				
Evening Phone:						
Email:		Date of Birth:	Gender:			
ICD10 Diagnosis Code:						
Chest Circumference:		Abdomen Measurement:				
(Measure fullest part of chest at nipple line)		(Measure largest circumference of abdomen at belly button line)				
Primary Insurance Provider:		Secondary Insurance Provider:				
BELOW THIS	LINE TO BE COMPLETED	BY A HEALTHCARE PRO	OVIDER ONLY			
Airway Clearance Therapy Tried and Faile	ed. This must be docu	mented in the natie	nt's progress notes			
Have alternative airway clearance techniques			nt 3 progress notes.			
Please indicate methods of airway clearance p						
CPT (manual or percussor)	Oscillating PEP (Flut		Pep Valve, Pep Mask)			
Huff coughing	☐ Breathing technique					
Hypertonic saline	Suctioning	(*Notes must document it is prescribed for secretion mobilization)				
2. Check all reasons why the above therapy failed	-	appropriate for this pat	ient:			
Cannot tolerate positioning/hand CPT			☐ Did not mobilize secretions ☐ Other			
Physical limitations of caregiver						
☐ Gastroesophageal reflux (GERD)	Severe arthritis, oste	eoporosis	Resistance to therapy			
Cognitive level	Unable to form mou	ıth seal	Artificial airway			
3. For Cystic Fibrosis or Neuromuscular patients, tl	he following must be doc	umented in the patient's	s progress notes. Please attach records with Rx.			
Documentation supporting diagnosis	Tried and failed a les	sser airway clearance te	chnique indicated above			
4. For Bronchiectasis patients, please check Yes o	or No to the following que	estion:				
Has there been a CT scan confirming Bronchie	ctasis diagnosis? 🔲 YES	S 🗌 NO If "Yes" please	e include copy of CT scan interpretation.			
In addition, the following medical history in th	e past year must be docu	mented in the patient's	s progress notes. Please attach records with Rx.			
=	infections, requiring anti	biotics in the last 12 mo	onths, documented at least two separate times			
OR ☐ Daily productive cough for more than s	iv continuous months					
Daily productive cough for more than s	ix continuous months					
Rx: High Frequency Chest Wall Oscill	ation (HFCWO HCF	PCS E0483)				
Start Date: Check need of Le	ngth: Lifetime (99)	Other				
Dispense one AffloVest by Tactile Medic	=		n/E0483			
Frequency of use (standard): Use the Af	floVest at 5Hz–20Hz for 3	0 minute treatments tw	vice per day (minimum of 15 minutes per day)			
Frequency of use (custom): Use the Affle	oVest at Hz	for minu	ute treatments per day			
Please check box if nebulizer therapy to	be used in conjunction	with HFCWO				
Physician Signature:			Date:			
Physician Printed Name:			NPI Number:			
Physician Address:						
City:	S	tate:	Zip:			
			il:			
Designated DME:						
I certify the accuracy of this Rx for the AffloVest Airway Clearance System a is true, accurate, and completed to the best of my knowledge. The patient the authorized AffloVest distributor by request. By providing this form to a information to process this order.	record contains the supplementary do	ocumentation to substantiate the me	edical necessity of the AffloVest and physician notes will be provided to			
**AffloVest requires a doctor's prescription for treatment by High Frequency Medicaid and private health insurance reimbursement under the Healthca Department of Veterans Affairs/Tricare. Patients must qualify to meet insur	re Common Procedure Coding Systen					
Durable Medical Equipment companies are ultimately responsible for en		a for a specific insurance plan and ۱	patient situation are satisfied.			

Tactile Medical

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Tactile

# **Medicare Approved ICD-10 Codes for AffloVest HFCWO Therapy (HCPCS E0483)**

### **Medicare Requirements for Bronchiectasis:**

1. Required: CT scan confirming diagnosis of bronchiectasis.

#### AND

2. Required: Daily productive cough for more than six continuous months.

#### OR

Frequent (i.e., more than two/year) exacerbations requiring antibiotic therapy in the last 12 months.

#### AND

3. Required: Documentation (chart notes) of another treatment tried to mobilize secretions and clearly indicating the other technique or device has failed.

ICD-10 CODE	DESCRIPTION
J47.0	Bronchiectasis with acute lower respiratory infection
J47.1	Bronchiectasis with (acute) exacerbation
J47.9	Bronchiectasis, uncomplicated
Q33.4	Congenital bronchiectasis

## Medicare Requirements for Other Respiratory, Cystic Fibrosis and Neuromuscular Conditions\*:

Physician's order that includes: AffloVest prescription, qualifying Dx, chart notes to support the Dx and well-documented failure of standard treatments to adequately mobilize retained secretions.

### **ICD-10 CODE/DESCRIPTION**

Dua abi		C71 021	Autocomo I domino et limbo etello escapio e dustro elec
Bronchie			Autosomal dominant limb girdle muscular dystrophy
J47.0	Bronchiectasis with acute lower respiratory infection	G/1.032	Autosomal recessive limb girdle muscular dystrophy due to
J47.1	Bronchiectasis with (acute) exacerbation	C71 022	calpain-3 dysfunction
J47.9	Bronchiectasis, uncomplicated		Limb girdle muscular dystrophy due to dysferlin dysfunction
Q33.4	Congenital bronchiectasis		Limb girdle muscular dystrophy due to sarcoglycan dysfunction, unspecified
Cystic Fi			Limb girdle muscular dystrophy due to alpha sarcoglycan dysfunction
E84.0	Cystic fibrosis with pulmonary manifestations		Limb girdle muscular dystrophy due to beta sarcoglycan dysfunction
E84.9	Cystic fibrosis, unspecified		Limb girdle muscular dystrophy due to other sarcoglycan dysfunction
Infection	ıs/Immune	G71.035	Limb girdle muscular dystrophy due to anoctamin-5 dysfunction
A15.0	Tuberculosis of lung	G71.036	Limb girdle muscular dystrophy due to fukutin related protein dysfunction Other limb girdle muscular dystrophy
Dolvion	3	G71.036 G71.039	Limb girdle muscular dystrophy, unspecified
Polyiom B91	Sequelae of poliomyelitis	G71.039 G71.09	Other specified muscular dystrophies
G14	Postpolio syndrome		•
	• •		s of the Diaphragm
	c/Metabolic Disorders	J98.6	Disorders of diaphragm
E74.02	Pompe disease	Myopath	ies
E74.05	Lysosome-associated membrane protein 2 [LAMP2] deficiency	G71.20	Congenital myopathy, unspecified
G71.11	Myotonic muscular dystrophy	G71.21	Nemaline myopathy
G71.12	Myotonia congenita	G71.220	X-linked myotubular myopathy
G71.13	Myotonic chondrodystrophy	G71.228	Other centronuclear myopathy
G71.14	Drug induced myotonia	G71.29	Other congenital myopathy
G71.19	Other specified myotonic disorders	G71.3	Mitochondrial myopathy, not elsewhere classified
Motor N	euron/Neuromuscular/Anterior Horn Cell Disease	G71.8	Other primary disorders of muscles
G12.0	Infantile spinal muscular atrophy, type I [Werdnig-Hoffman]	G72.0	Drug-induced myopathy
G12.1	Other inherited spinal muscular atrophy	G72.1	Alcoholic myopathy
G12.20	Motor neuron disease, unspecified	G72.2	Myopathy due to other toxic agents
G12.21	Amyotrophic lateral sclerosis	G72.41	Inclusion body myositis [IBM]
G12.22	Progressive bulbar palsy	G72.49	Other inflammatory and immune myopathies, not elsewhere classified
G12.23	Primary lateral sclerosis	G72.89	Other specified myopathies
G12.24	Familial motor neuron disease	G72.9	Myopathy, unspecified
G12.25	Progressive spinal muscle atrophy	G73.7	Myopathy in diseases classified elsewhere
G12.29	Other motor neuron disease	M33.02	Juvenile dermatomyositis with myopathy
G12.8	Other spinal muscular atrophies and related syndromes	M33.12	Other dermatomyositis with myopathy
G12.9	Spinal muscular atrophy, unspecified	M33.22	Polymyositis with myopathy
Multiple	Sclerosis	M33.92	Dermatopolymyositis, unspecified with myopathy
G35.A	Relapsing remitting MS	M34.82	Systemic sclerosis with myopathy
G35.B0	Primary progressive MS	M35.03	Sicca syndrome with myopathy
G35.B1	Active primary progressive MS	Quadriple	egia
G35.B2	Non-active primary progressive MS	G80.0	Spastic quadriplegic cerebral palsy
G35.C0	Secondary progressive MS	G82.50	Quadriplegia, unspecified
G35.C1	Active secondary progressive MS	G82.51	Quadriplegia, C1-C4 complete
G35.C2	Non-active secondary progressive MS	G82.52	Quadriplegia, C1-C4 incomplete
G35.D	Multiple sclerosis, unspecified	G82.53	Quadriplegia, C5-C7 complete
Muscula	r Dystrophy	G82.54	Quadriplegia, C5-C7 incomplete
G71.00	Muscular dystrophy, unspecified	Myastha	nia Gravis
G71.01	Duchenne or Becker muscular dystrophy	G70.00	Myasthenia gravis without acute exacerbation
G71.01	Fascioscapulohumeral muscular dystrophy	G70.00 G70.01	Myasthenia gravis with acute exacerbation  Myasthenia gravis with acute exacerbation
27		0.0.01	Myastrema gravis with acute exacerbation

\*cms.gov/medicare-coverage-database/view/lcd.aspx?LCDId=33785&ContrlD=140



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