




ICD-10 Codes


Medicare-Covered Diagnoses for HFCWO

To qualify for reimbursement, a bronchiectasis patient must have the following documented:

- 

HRCT Scan
Confirming bronchiectasis diagnosis
- 

Daily Productive Cough
For at least 6 consecutive months
- OR**
- 

Frequent Exacerbations Requiring Antibiotic Therapy
(3 or more annually)
- 

Failure of Standardized Treatments
Such as OPEP or CPT to mobilize retained secretions

Bronchiectasis

J47.0 Bronchiectasis with acute lower respiratory infection

J47.1 Bronchiectasis with (acute) exacerbation

J47.9 Bronchiectasis, uncomplicated

Q33.4 Congenital bronchiectasis

Cystic Fibrosis

E84.0 Cystic fibrosis with pulmonary manifestations

E84.9 Cystic fibrosis, unspecified

Late Effects of Poliomyelitis

B91 Sequelae of poliomyelitis

G14 Postpolio syndrome

Disorders of Diaphragm

J98.6 Disorders of Diaphragm

Infectious/Immune

A15.0 Tuberculosis of lung

"Amazing for bronchiectasis, I've seen fewer pneumonias"

— Latecia, SmartVest User



Motor Neuron/Spinal Muscular Atrophy

- G12.0** Infantile spinal muscular atrophy, type I [Werdnig Hoffman]
- G12.1** Other inherited spinal muscular atrophy
- G12.20** Motor neuron disease, unspecified
- G12.21** Amyotrophic lateral sclerosis
- G12.22** Progressive bulbar palsy
- G12.23** Primary lateral sclerosis
- G12.24** Familial motor neuron disease
- G12.25** Progressive spinal muscle atrophy
- G12.29** Other motor neuron disease
- G12.8** Other spinal muscular atrophies and related syndromes
- G12.9** Spinal muscular atrophy, unspecified

Multiple Sclerosis

- G35.A** Relapsing-remitting multiple sclerosis
- G35.B0** Primary progressive multiple sclerosis, unspecified
- G35.B1** Active primary progressive multiple sclerosis
- G35.B2** Non-active primary progressive multiple sclerosis
- G35.C0** Secondary progressive multiple sclerosis, unspecified
- G35.C1** Active secondary progressive multiple sclerosis
- G35.C2** Non-active secondary progressive multiple sclerosis
- G35.D** Multiple sclerosis, unspecified

Myasthenia Gravis

- G70.00** Myasthenia gravis without (acute) exacerbation
- G70.01** Myasthenia gravis with (acute) exacerbation

Muscular Dystrophy

- G71.00** Muscular dystrophy, unspecified
- G71.01** Duchenne or Becker muscular dystrophy
- G71.02** Fascioscapulohumeral muscular dystrophy
- G71.031** Autosomal dominant limb girdle muscular dystrophy
- G71.032** Autosomal recessive limb girdle muscular dystrophy due to calpain-3 dysfunction
- G71.033** Limb girdle muscular dystrophy due to dysferlin dysfunction
- G71.0340** Limb girdle muscular dystrophy due to sarcoglycan dysfunction, unspecified
- G71.0341** Limb girdle muscular dystrophy due to alpha sarcoglycan dysfunction
- G71.0342** Limb girdle muscular dystrophy due to beta sarcoglycan dysfunction
- G71.0349** Limb girdle muscular dystrophy due to other sarcoglycan dysfunction
- G71.035** Limb girdle muscular dystrophy due to anoctamin-5 dysfunction
- G71.036** Limb girdle muscular dystrophy due to fukutin related protein dysfunction

- G71.038** Other limb girdle muscular dystrophy
- G71.039** Limb girdle muscular dystrophy, unspecified
- G71.09** Other specified muscular dystrophies
- G71.11** Myotonic muscular dystrophy

Myotonic and Metabolic Disorders

- E74.02** Pompe disease
- E74.05** Lysosome-associated membrane protein 2 [LAMP2] deficiency
- G71.12** Myotonia congenita
- G71.13** Myotonic chondrodystrophy
- G71.14** Drug induced myotonia
- G71.19** Other specified myotonic disorders

Other Myopathies

- G71.20** Congenital myopathies, unspecified
- G71.21** Nemaline myopathy
- G71.220** X-linked myotubular myopathy
- G71.228** Other centronuclear myopathy
- G71.29** Other congenital myopathy
- G71.3** Mitochondrial myopathy, not elsewhere classified
- G71.8** Other primary disorders of muscles
- G72.0** Drug-induced myopathy
- G72.1** Alcoholic myopathy
- G72.2** Myopathy due to other toxic agents
- G72.41** Inclusion body myositis [IBM]
- G72.49** Other inflammatory and immune myopathies, not elsewhere classified
- G72.89** Other specified myopathies
- G72.9** Myopathy, unspecified
- G73.7** Myopathy in diseases classified elsewhere
- M33.02** Other congenital myopathy
- M33.12** Other dermatomyositis with myopathy
- M33.22** Polymyositis with myopathy
- M33.92** Dermatopolymyositis, unspecified with myopathy
- M34.82** Systemic sclerosis with myopathy
- M35.03** Sicca syndrome with myopathy

Quadriplegia

- G80.0** Spastic quadriplegic cerebral palsy
- G82.50** Quadriplegia, unspecified
- G82.51** Quadriplegia, C1-C4 complete
- G82.52** Quadriplegia, C1-C4 incomplete
- G82.53** Quadriplegia, C5-C7 complete
- G82.54** Quadriplegia, C5-C7 incomplete

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