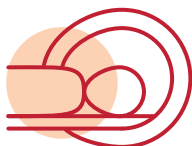


Medicare-Covered Diagnoses

ICD-10 Codes Covered for HFCWO

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

1



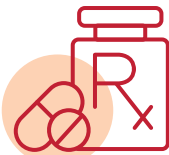
HRCT scan
confirming
bronchiectasis
diagnosis

2



**Daily productive
cough**
for at least
six continuous months

OR



**Frequent
exacerbations
requiring antibiotic
therapy** (i.e. more
than twice annually)

3



**Failure of standard
treatments**
such as OPEP
to mobilize
retained secretions

Bronchiectasis Diagnoses Covered by Medicare

Bronchiectasis

- J47.0** Bronchiectasis with acute lower respiratory infection
- J47.1** Bronchiectasis with (acute) exacerbation

- J47.9** Bronchiectasis, uncomplicated
- Q33.4** Congenital bronchiectasis

Other Diagnoses Covered by Medicare

Cystic Fibrosis

- E84.0** Cystic fibrosis with pulmonary manifestations
- E84.9** Cystic fibrosis, unspecified

Infectious / Immune

- A15.0** Tuberculosis of lung
- D84.1** Defects in the complement system

Disorders of Diaphragm

- J98.6** Disorders of diaphragm

Late Effects of Poliomyelitis

- B91** Sequelae of poliomyelitis
- G14** Postpolio syndrome

continues other side...

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Motor Neuron / Neuromuscular / Anterior Horn Cell Disease

- 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** Multiple sclerosis

Muscular Dystrophy

- G71.00** Muscular dystrophy, unspecified
- G71.01** Duchenne or Becker muscular dystrophy
- G71.02** Fascioscapulohumeral muscular dystrophy
- G71.09** Other specified muscular dystrophies

Myotonic and Metabolic Disorders

- D81.810** Biotinidase deficiency
- G71.11** Myotonic muscular dystrophy
- G71.12** Myotonia congenita
- G71.13** Myotonic chondrodystrophy
- G71.14** Drug induced myotonia
- G71.19** Other specified myotonic disorders

Other Myopathies

- G71.2** Congenital myopathies
- 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
- G72.89** Other specified myopathies
- G73.7** Myopathy in diseases classified elsewhere
- M33.02** Juvenile dermatomyositis with 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

- 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

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

— Latecia, SmartVest user



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EM00541.2019-09