

Autosomalna recesywna choroba Charcota, Mariego i Tootha typu 2X

Kod Orpha: 466775 Kod OMIM: 616668

Opis choroby *

Definicja

A rare autosomal recessive axonal hereditary motor and sensory neuropathy characterized by childhood to adult onset of slowly progressive, sometimes asymmetric distal muscle weakness and atrophy, as well as sensory impairment, predominantly of the lower limbs. Additional common features include pes cavus, kyphoscoliosis, ankle contractures, tremor, or urogenital dysfunction. Fasciculations and proximal involvement may be seen in some cases. Patients usually remain ambulatory.

Dane

Klasyfikacja	Synonimy
Choroba	ARCMT2X ARCMT2X Autosomal recessive Charcot-Marie-Tooth disease type 2 due to SPG11 mutation CMT2X
	Autosomal recessive Charcota, Mariego i Tootha typu 2 spowodowana SPG11
	CMT2X
	Autosomal recessive Charcot-Marie-Tooth disease type 2 due to SPG11 mutation CMT2X

Kod ORPHA	Kod OMIM	Kod ICD10
466775	616668	G60.0

Kod ICD11

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*Źródło

orphanet

Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.

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