

Choroba Charcota, Mariego i Tootha typu 1C

Kod Orpha: 101083 Kod OMIM: 601098

Opis choroby *

Definicja

A rare, autosomal dominant, hereditary, demyelinating motor and sensory neuropathy which may present either as a classic Charcot-Marie-Tooth disease phenotype with distal motor weakness and wasting, gait difficulties, paresthesias, decreased vibration and pain sensation, or as a milder, predominantly sensory form with transient paresthesias, decreased sensation and distal pain in upper or lower limbs, without significant motor weakness. Pes cavus is a common feature, and additional symptoms may include hand tremor and decreased or absent deep tendon reflexes.

Dane

Klasyfikacja

Choroba

Synonimy

CMT1C

CMT1C

Kod ORPHA

101083

Kod OMIM

601098

Kod ICD10

G60.0

Kod ICD11

8C20.0

*[Źródło](#)

orphanet

Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.