Opis choroby *

Definicja

A rare hereditary motor and sensory neuropathy disorder characterized by the typical CMT phenotype (slowly progressive distal muscle atrophy and weakness in upper and lower limbs, distal sensory loss in extremities, reduced or absent deep tendon reflexes and foot deformities) with nerve biopsy demonstrating demyelinating and axonal changes and nerve conduction velocities varying from the demyelinating to axonal range.

Dane

Klasyfikacja Choroba Synonimy CMTDIF CMTDIF

Kod ORPHA

352670

Kod OMIM 615185

Kod ICD10 G60.0

Kod ICD11 8C20.2

*Źródło

orphanet