

Choroba Charcota, Mariego i Tootha typu 1E

Kod Orpha: 90658 Kod OMIM: 118300

Opis choroby *

Definicja

A rare subtype of CMT1 characterized by a variable clinical presentation. Onset within the first two years of life with a delay in walking is not uncommon; however, onset may occur later. CMT1E is caused by point mutations in the *PMP22* (17p12) gene. The disease severity depends on the particular *PMP22* mutation, with some cases being very mild and even resembling hereditary neuropathy with liability to pressure palsies, while others having an earlier onset with a more severe phenotype (reminiscent of Dejerine-Sottas syndrome) than that seen in CMT1A, caused by gene duplication. These severe cases may also report deafness and much slower motor nerve conduction velocities compared to CMT1A patients.

Dane

Klasyfikacja	Synonimy
Choroba	CMT1E
	Choroba Charcota, Mariego i Tootha - głuchota
	CMT1E
	Charcot-Marie-Tooth disease-deafness syndrome
	Charcot-Marie-Tooth disease-hearing loss syndrome

Kod ORPHA	Kod OMIM	Kod ICD10
90658	118300	G60.0

Kod ICD11
8C20.0

*[Źródło](#)

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Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.

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