

Opis choroby *

Definicja

A severe form of axonal Charcot-Marie-Tooth disease, a peripheral sensorimotor neuropathy, with onset in the 2nd or 3rd decade, characterized by ulcerations and infections of feet. Symmetric and distal weakness develops mostly in the legs together with a severe symmetric distal sensory loss, tendon reflexes are only reduced at ankles and foot deformities, including pes cavus or planus and hammer toes, appear in childhood.

Dane

Klasyfikacja

Choroba

Synonimy

CMT2B

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Kod ORPHA

99936

Kod OMIM

600882

Kod ICD10

G60.0

Kod ICD11

8C20.1

*Źródło

orphanet