

Opis choroby *

Definicja

A complex form of hereditary spastic paraplegia characterized by spastic paraplegia, demyelinating peripheral sensorimotor neuropathy, poikiloderma (manifesting with loss of eyebrows and eyelashes in childhood in addition to delicate, smooth, and wasted skin) and distal amyotrophy (presenting after puberty). There have been no further descriptions in the literature since 1992.

Dane

Klasyfikacja

Choroba

Synonimy

Antinolo-Nieto-Borrego syndrome

Zespół Antinolo, Nieto i Borrego

Kod ORPHA

2821

Kod OMIM

182815

Kod ICD10

G11.4

Kod ICD11

-

*Źródło

orphanet