Opis choroby *

Definicja

A rare genetic leukodystrophy identified in families of Ashkenazi Jewish descent, characterized by infancy onset of severe global developmental delay with very limited or absent speech and sometimes complete absence of motor development, hypotonia, spasticity, and acquired microcephaly. Seizures, hearing loss, visual impairment, and autonomic dysfunction have also been described. Brain imaging shows delayed myelination and other white matter abnormalities.

Dane

Klasyfikacja Synonimy

Choroba VPS11-related autosomal recessive

hypomyelinating leukoencephalopathy Autosomalna recesywna leukoencefalopatia

hipomielinizacyjna zależna od VPS11

Kod ORPHA 466934

Kod OMIM 616683

Kod ICD10

G93.8

Kod ICD11

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*Źródło

orphanet