Opis choroby *

Definicja

Epileptic encephalopathy with global cerebral demyelination is a rare mitochondrial substrate carrier disorder characterized by severe muscular hypotonia, seizures (with or without episodic apnea) beginning in the first year of life, and arrested psychomotor development (affecting mainly motor skills). Severe spasticity with hyperreflexia has also been reported. Global cerebral hypomyelination is a characteristic imaging feature of this disease.

Dane

Klasyfikacja Synonimy

Choroba AGC1 deficiency

Niedobór mitochondialnego nośnika 1

asparaginianu-glutaminianu

Mitochondrial aspartate-glutamate carrier 1

deficiency

Kod ORPHA Kod OMIM Kod ICD10

353217 612949 E88.8

Kod ICD11

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*Źródło

orphanet