## Opis choroby \*

Definicja

A rare genetic neurometabolic disease characterized by early neonatal refractory seizures, hypotonia, and respiratory failure. Brain imaging reveals simplified gyral pattern of the frontal lobes, white matter abnormalities, gliosis and volume loss in various brain regions, and vasogenic edema. Serum glutamine levels are significantly elevated. Death occurs within weeks after birth.

Dane

Klasyfikacja

Choroba

Kod ORPHA 557064

**Kod OMIM** 618328

Kod ICD10 E88.8

**Kod ICD11** 

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## \*Źródło

orphanet