

Opis choroby *

Definicja

A rare congenital disorder of glycosylation characterized by neonatal onset of global developmental delay, hypotonia, failure to thrive, hematological/immunological abnormalities, recurrent infections, liver involvement (with hepatosplenomegaly, cholestasis, fibrosis, or cirrhosis), and enteropathy. Additional reported manifestations include dysmorphic craniofacial features (such as microcephaly, broad palpebral fissures, and retrognathia), hypohidrosis, hyperkeratosis, and cardiac and musculoskeletal anomalies. Brain imaging may show hypoplastic corpus callosum, cerebral and cerebellar atrophy, and enlarged ventricles.

Dane

Klasyfikacja

Choroba

Synonimy

CDG syndrome type IIL

Wrodzone zaburzenia glikozylacji typu 2I

Wrodzone zaburzenia glikozylacji typu III

CDG-IIL

CDG2L

Congenital disorder of glycosylation type 2I

Congenital disorder of glycosylation type IIL

Kod ORPHA

464443

Kod OMIM

614576

Kod ICD10

E77.8

Kod ICD11

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*Źródło

orphanet