

## Opis choroby \*

### Definicja

A rare genetic neurological disorder characterized by subacute encephalopathy with confusion, seizures, and movement disorder, often following a history of febrile illness. Imaging may reveal bilateral lesions in the basal ganglia. The disease usually becomes symptomatic in childhood and is life-threatening if left untreated, but symptoms can be reversed and progression prevented by treatment with high doses of biotin and thiamine.

### Dane

#### Klasyfikacja

Choroba

#### Synonimy

BBGD

BBGD

BTBGD

Choroba zwojów podstawnych wrażliwa na biotyne-tiaminę

BTBGD

Biotin-responsive basal ganglia disease

#### Kod ORPHA

65284

#### Kod OMIM

607483

#### Kod ICD10

G93.8

#### Kod ICD11

5C63.Y

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#### [\\*Źródło](#)

orphanet