

# Acyduria D-glicerynowa

## Kod Orpha: 941 Kod OMIM: 220120

### Opis choroby \*

#### Definicja

A rare inborn error of metabolism characterized by abnormal urinary excretion of D-glyceric acid due to D-glycerate kinase deficiency. Reported manifestations are highly variable and include a severe encephalopathic picture, chronic metabolic acidosis, developmental delay, intellectual disability, microcephaly, seizures, behavioral abnormalities, as well as only mild speech delay and apparently normal development.

#### Dane

Klasyfikacja	Synonimy
Choroba	D-glycerate kinase deficiency
	Kwasica D-glicerynowa
	Niedobór kinazy D-glicerynowej
	D-glyceric acidemia
<b>Kod ORPHA</b>	<b>Kod OMIM</b>
941	220120
<b>Kod ICD11</b>	<b>Kod ICD10</b>
5C50.7Y	E74.8

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\*Źródło

orphanet

### Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.