

# Niedobór dehydrogenazy glukozo-6-fosfatazy klasy I

Kod Orpha: 466026 Kod OMIM: 300908

## Opis choroby \*

### Definicja

A rare constitutional hemolytic anemia due to an enzyme disorder characterized by severe glucose-6-phosphate dehydrogenase deficiency (typically <10% residual enzyme activity) associated with chronic non-spherocytic hemolytic anemia of highly variable severity. Patients are at risk of developing neonatal jaundice (potentially leading to kernicterus), gallstones, and reticulocytosis and splenomegaly. They have an increased susceptibility to oxidizing agents provoking episodes of acute hemolysis. Favism, which describes the occurrence of an acute hemolytic reaction in response to the ingestion of fava beans, is more common in infants and young children.

### Dane

#### Klasyfikacja

Choroba

#### Synonimy

Class I G6PD deficiency  
Ciężka niedokrwistość hemolityczna  
spowodowana niedoborem G6PD  
Niedobór G6PD klasy I  
Severe hemolytic anemia due to G6PD deficiency

#### Kod ORPHA

466026

#### Kod OMIM

300908

#### Kod ICD10

D55.0

#### Kod ICD11

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\*[Źródło](#)

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## **Rozszerzony opis choroby**

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.

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