

# Niedokrwistość sierpowata - beta-talasemia

## Kod Orpha: 251359 Kod OMIM:

### Opis choroby \*

#### Definicja

A rare, genetic hemoglobinopathy that affects red blood cells both in the production of abnormal hemoglobin, as well as the decreased synthesis of beta globin chains. Clinical manifestations depend on the amount of residual beta globin chains production, and are similar to sickle cell disease, including anemia, vascular occlusion and its complications, acute episodes of pain, acute chest syndrome, pulmonary hypertension, sepsis, ischemic brain injury, splenic sequestration crisis and splenomegaly.

#### Dane

Klasyfikacja	Synonimy
Choroba	HbS-beta-thalassemia syndrome HbS - beta-talasemia

Kod ORPHA	Kod OMIM	Kod ICD10
251359	-	D57.2

Kod ICD11  
3A51.4

---

\*Źródło

orphanet

### Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.