

# **Wrodzona atransferynemia**

## **Kod Orpha: 1195 Kod OMIM: 209300**

### **Opis choroby \***

#### **Definicja**

Congenital atransferrinemia is a very rare hematologic disease caused by a transferrin (TF) deficiency and characterized by microcytic, hypochromic anemia (manifesting with pallor, fatigue and growth retardation) and iron overload, and that can be fatal if left untreated.

#### **Dane**

##### **Klasyfikacja**

Choroba

##### **Synonimy**

Congenital hypotransferrinemia

Wrodzona hipotransferynemia

##### **Kod ORPHA**

1195

##### **Kod OMIM**

209300

##### **Kod ICD10**

E88.0

##### **Kod ICD11**

5D0Y

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\*Źródło

orphanet

### **Rozszerzony opis choroby**

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.