

# Rodzinny zespół chylomikronemii

## Kod Orpha: 444490 Kod OMIM: 615947

### Opis choroby \*

#### Definicja

A rare genetic hyperlipidemia characterized by excessive increase in plasma triglyceride levels due to the accumulation of chylomicrons. Clinical manifestations include recurrent episodes of severe acute pancreatitis, abdominal pain, nausea, fatigue, diarrhea, constipation, hepatosplenomegaly, eruptive xanthomas, and failure to thrive. Children may often be asymptomatic. The condition is not associated with severe atherosclerosis.

#### Dane

#### Klasyfikacja

Choroba

Kod ORPHA  
444490

Kod OMIM  
615947

Kod ICD10  
E78.3

#### Kod ICD11

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[\\*Źródło](#)

orphanet

### Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.