

# **Wrodzona biegunka spowodowana niedoborem komórek enteroendokrynowych**

## **Kod Orpha: 83620 Kod OMIM: 610370**

### **Opis choroby \***

#### **Definicja**

A very rare genetic gastroenterological disease characterized by severe malabsorptive diarrhea (requiring parenteral nutrition and disappearing at fasting) due to a lack of intestinal enteroendocrine cells. It is associated with early-onset (within the first weeks of life) dehydration, metabolic acidosis and diabetes mellitus (that can develop until late childhood). Patient may display various degrees of pancreatic insufficiency that does not explain diarrhea, as it is not reduced with pancreatic enzyme supplementation. Central hypogonadism (developing in the second decade), as well as an association with celiac disease have been reported.

#### **Dane**

##### **Klasyfikacja**

Choroba

##### **Synonimy**

Congenital malabsorptive diarrhea due to paucity of enteroendocrine cells  
Anedokrynoza jelitowa

##### **Kod ORPHA**

83620

##### **Kod OMIM**

610370

##### **Kod ICD10**

P78.3

##### **Kod ICD11**

DA90.Y

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\*Źródło

orphanet

### **Rozszerzony opis choroby**

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.

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