

Opis choroby *

Definicja

A rare pancreatic disease characterized by a most commonly single, unilocular, thin-walled cystic lesion which may be located anywhere within the pancreas (but is more frequently found in the body and tail) and does not communicate with the pancreatic ductal system. Patients may be asymptomatic or present with signs and symptoms of gastrointestinal or biliary obstruction, or pancreatitis. The condition can be isolated or occur in association with other anomalies (such as von Hippel-Lindau disease or polycystic kidney disease).

Dane

Klasyfikacja

Wada morfologiczna

Synonimy

Neonatal congenital pancreatic cyst
Noworodkowa wrodzona torbiel trzustki
Prawdziwa wrodzona torbiel trzustki
True congenital pancreatic cyst

Kod ORPHA

313906

Kod OMIM

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Kod ICD10

Q45.2

Kod ICD11

LB21.Y

*Źródło

orphanet