

Nerka dysplastyczna wielotorbielowata

Kod Orpha: 1851 Kod OMIM:

Opis choroby *

Definicja

A rare congenital anomaly of the kidney and urinary tract (CAKUT) in which one or both kidneys (unilateral or bilateral MCDK respectively) are large, distended by multiple cysts, and non-functional. Unilateral MCDK is typically asymptomatic if the other kidney is fully functional but may occasionally present with abdominal obstructive signs when the cysts become too large. Bilateral MCDK is considered a lethal entity and neonates present with features of the Potter sequence, severe pulmonary hypoplasia and severe renal failure, and generally die shortly after birth.

Dane

Klasyfikacja

Wada morfologiczna

Synonimy

MCDK
Dysplazja wielotorbielowata nerki
MCDK
Multicystic renal dysplasia

Kod ORPHA

1851

Kod OMIM

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Kod ICD10

Q61.4

Kod ICD11

LB30.9

[*Źródło](#)

orphanet

Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.

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