

Rodzinna glikozuria nerkowa

Kod Orpha: 69076 Kod OMIM: 233100

Opis choroby *

Definicja

A rare, genetic, glucose transport disorder characterized by the presence of persistent isolated glucosuria in the absence of both proximal tubular dysfunction and hyperglycemia. The disorder is benign in the majority of cases although it may occasionally manifest with polyuria, enuresis, a mild growth and pubertal maturation delay, hypercalciuria, aminoaciduria and, in severe cases, increased incidence of urinary infections and episodic dehydration and ketosis during pregnancy and starvation.

Dane

Klasyfikacja	Synonimy
Choroba	Familial renal glycosuria Niedobór SGLT2 SGLT2 deficiency
Kod ORPHA	Kod OMIM
69076	233100
Kod ICD11	Kod ICD10
5C61.5	E74.8

*Źródło

orphanet

Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.