## Opis choroby \*

## Definicja

A rare, congenital, isolated hyperinsulinism disorder characterized by neonatal presentation of severe refractory hypoglycemia in the first two days of life, with limited response to medical management, sometimes requiring pancreatic resection. Newborns are often large for gestational age with mild to moderate hepatomegaly and diffuse form of hyperinsulinism due to SUR1 deficiency. Persistent hypoglycemia, hyperglycemia and type1 diabetes mellitus may develop later in life. Life-threatening hypoglycemic coma or status epilepticus have also been associated.

Dane

**Klasyfikacja** Synonimy

Choroba Autosomal recessive hyperinsulinemic

hypoglycemia due to SUR1 deficiency Autosomalna recesywna hipoglikemia

hiperinsulinemiczna z powodu niedoboru SUR1

**Kod ICD10** 

Kod ORPHA Kod OMIM

79643 256450 E16.1

**Kod ICD11** 

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## <u>\*Źródło</u>

orphanet