

Opis choroby *

Definicja

A rare, genetic, mitochondrial oxidative phosphorylation disorder characterized by a potentially life-threatening, severe myopathy manifesting in the neonatal to early infantile period, followed by marked, spontaneous improvement of muscular function by early childhood. Associated biochemical findings include lactic acidosis and a transient, marked decrease in respiratory chain activity.

Dane

Klasyfikacja

Choroba

Synonimy

Benign COX deficiency
Dziecięcy odwracalny Niedobór oksydazy cytochromu c
Łagodny Niedobór COX
Miopatia mitochondrialna z odwracalnym niedoborem COX
Miopatia mitochondrialna z odwracalnym niedoborem kompleksu IV
Infantile reversible cytochrome C oxidase deficiency myopathy
Mitochondrial myopathy with reversible COX deficiency
Mitochondrial myopathy with reversible complex IV deficiency
Reversible infantile cytochrome C oxidase deficiency
Reversible infantile respiratory chain deficiency

Kod ORPHA

254864

Kod OMIM

500009

Kod ICD10

G71.3

Kod ICD11

8C73.Y

[*Źródło](#)

orpho:net