

Miopatia mitochondrialna dziedziczona od matki

Kod Orpha: 254788 Kod OMIM:

Opis choroby *

Definicja

A group of rare mitochondrial oxidative phosphorylation disorders due to mitochondrial DNA anomalies characterized by progressive, most commonly proximal, myopathy with variable degrees of weakness, exercise-induced muscle pain, and fatigue. Progressive external ophthalmoplegia is often observed. Additional features include neurological signs and symptoms (such as seizures, stroke-like episodes, or developmental delay), cardiomyopathy, involvement of liver, kidneys, and gastrointestinal tract, and diabetes. Lactic acidosis is frequently present, while recurrent rhabdomyolysis and myoglobinuria are rare. Muscle biopsy may reveal the presence of ragged-red fibers and a mosaic pattern of cytochrome c oxidase-negative fibers.

Dane

Klasyfikacja

Grupa fenomenów

Synonimy

Maternally-inherited mitochondrial myopathy
mtDNA-related mitochondrial myopathy

Kod ORPHA

254788

Kod OMIM

-

Kod ICD10

G71.3

Kod ICD11

-

[*Źródło](#)

orphanet

Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.

Orphanet - internetowa baza danych dotyczących rzadkich chorób i sierochych leków. ©INSERM 1999 -
Dostępna na stronie www.orphanet.pl