

Złożony defekt fosforylacji oksydacyjnej typu 7

Kod Orpha: 254930 Kod OMIM: 613559

Opis choroby *

Definicja

Combined oxidative phosphorylation defect type 7 is a rare mitochondrial disease due to a defect in mitochondrial protein synthesis characterized by a variable phenotype that includes onset in infancy or early childhood of failure to thrive and psychomotor regression (after initial normal development), as well as ocular manifestations (such as ptosis, nystagmus, optic atrophy, ophthalmoplegia and reduced vision). Additional manifestations include bulbar paresis with facial weakness, hypotonia, difficulty chewing, dysphagia, mild dysarthria, ataxia, global muscle atrophy, and areflexia. It has a relatively slow disease progression with patients often living into the third decade of life.

Dane

Klasyfikacja	Synonimy
Choroba	COXPD7
	COXPD7
	Severe C12ORF65-related COXPD
	Severe C12ORF65-related combined oxidative phosphorylation defect

Kod ORPHA	Kod OMIM	Kod ICD10
254930	613559	E88.8

Kod ICD11
5C53.23

[* Źródło](#)

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Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.

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