

Pasażowalna encefalopatia gąbczasta

Kod Orpha: 56970 Kod OMIM:

Opis choroby *

Definicja

A group of rare neurodegenerative diseases characterized by the accumulation of prions, abnormal variants of the cellular prion protein, primarily in brain tissue of affected individuals, as well as massive, rapid neuronal death, and an invariably fatal course. Human prion diseases most often occur sporadically but may also be of genetic origin or infectiously acquired. Irrespective of etiology, they are transmissible to other individuals.

Dane

Klasyfikacja	Synonimy
Kategoria	TSE
	Choroba prionowa
	Transmissible spongiform encephalopathy

Kod ORPHA	Kod OMIM	Kod ICD10
56970	-	-

Kod ICD11

*Źródło

orphanet

Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.