Opis choroby *

Definicja

A rare genetic neurological disorder characterized by infantile or childhood onset of recurrent acute encephalopathic episodes with cerebellar and extrapyramidal involvement following febrile illnesses. During the episodes, patients typically show sudden onset of truncal ataxia, occasionally accompanied by lethargy and impairment of speech, as well as choreic and athetoid movements, seizures, loss of deep tendon reflexes, and presence of pathological reflexes. Episodes last from day to weeks and may leave residual symptoms such as speech impairment and poor coordination. There have been no further descriptions in the literature since 1983.

Dane

Klasyfikacja Synonimy

Zespół wad wrodzonych Recurrent encephalophathy of childhood

Nawracająca encefalopatia dziecięca

 Kod ORPHA
 Kod OMIM
 Kod ICD10

 2672
 130950
 G93.4

Kod ICD11

_

*Źródło

orphanet