Opis choroby *

Definicja

A heterogenous group of cerebral cortical malformations characterized by excessive cortical folding and abnormal cortical layering that, depending on its topographic distribution, presents with variable combinations of neurological symptoms of varying severity such as epilepsy, developmental delay, intellectual disability, motor dysfunction (e.g. spasticity), and pseudobulbar palsy.

Dane

Klasyfikacja Grupa fenomenów

Kod ORPHA 35981 Kod OMIM

Kod ICD10 Q04.3

Kod ICD11 LA05.50

<u>*Źródło</u>

orphanet