

Opis choroby *

Definicja

A rare genetic developmental and epileptic encephalopathy (DEE) characterized by developmental delay, generalized epilepsy consisting of eyelid myoclonia with absences and myoclonic-atonic seizures, intellectual disability and autism spectrum disorder (ASD).

Dane

Klasyfikacja	Synonimy
Choroba	SYNGAP1-related DEE SYNGAP1-related DEE
Kod ORPHA 544254	Kod OMIM -

Kod ORPHA 544254	Kod OMIM -	Kod ICD10 G40.4
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Kod ICD11

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*Źródło

orphanet