Opis choroby *

Definicja

A rare genetic hyperkinetic movement disorder characterized predominantly by chorea of variable severity, associated with bilateral striatal abnormalities on cerebral MRI. The disease is scarcely progressive, and cognitive performance is preserved in the majority of cases, although mild cognitive delay has also been reported.

Dane

Klasyfikacja

Choroba

Kod ORPHA 494541

Kod OMIM 616922

Kod ICD10 G25.5

Kod ICD11

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*Źródło

orphanet