Opis choroby *

Definicja

A rare genetic disease characterized by the triad of adult-onset moderate to severe bilateral sensorineural hearing loss, premature graying of scalp hair, and essential tremor manifesting as involuntary shaking of the head. Additional pigmentation abnormalities have not been reported in this syndrome.

Dane

Klasyfikacja Synonimy

Choroba Sensorineural deafness-early graying-essential

tremor syndrome

Sensorineural deafness-early graying-essential

tremor syndrome

Kod ORPHA

66633

Kod OMIM

Kod ICD10

-

H90.3

Kod ICD11 LD2H.Y

*Źródło

orphanet