

Zespół Ushera typu 2

Kod Orpha: 231178 Kod OMIM: 611383

Opis choroby *

Definicja

A rare ciliopathy characterized by congenital moderate-to-severe deafness, retinitis pigmentosa developing in the first or second decade, and normal vestibular function. Congenital bilateral sensorineural hearing loss is mild to moderate in the low frequencies and severe to profound in the higher frequencies. Additional manifestations include night blindness, constricted visual field (tunnel vision), and later on decreased visual acuity sometimes ending with bare light perception.

Dane

Klasyfikacja

Podtyp kliniczny

Synonimy

USH2

USH2

Kod ORPHA

231178

Kod OMIM

611383

Kod ICD10

H35.5

Kod ICD11

LD2H.4

*[Źródło](#)

[orphanet](#)

Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.