

Opis choroby *

Definicja

A rare multiple congenital anomalies/dysmorphic syndrome characterized by anomalous ossification and skeletal patterning of the axial and appendicular skeleton, facial dysmorphism and conductive and sensorineural hearing loss.

Dane

Klasyfikacja

Choroba

Kod ORPHA
1826

Kod OMIM
617137

Kod ICD10
Q78.5

Kod ICD11
LD25.1

*[Źródło](#)

[orphanet](#)