Opis choroby *

Definicja

A rare, genetic, non-syndromic developmental defect of the eye disorder, with high clinical and genetic heterogeneity, most frequently characterized by bilateral, symmetrical, non-progressive cataracts which present at birth or in early-childhood. Additional ocular manifestations (e.g. anterior segment dysgenesis, colobomas, nystagmus, microcornea, microphthalmia, myopia) may be associated, however other organs/systems are usually not affected.

Dane

Klasyfikacja Choroba

Kod ORPHA 91492

Kod OMIM 601885 Kod ICD10 Q12.0

Kod ICD11 LA12.1

<u>*Źródło</u>

orphanet