

Opis choroby *

Definicja

A rare, genetic, lens position anomaly disease characterized by bilateral congenital blepharoptosis, ectopia lentis and high grade myopia. Additional reported manifestations include abnormally long eye globes and signs of levator aponeurosis disinsertion. There have been no further descriptions in the literature since 1982.

Dane

Klasyfikacja

Choroba

Kod ORPHA

1259

Kod OMIM

110150

Kod ICD10

Q15.8

Kod ICD11

LA14.0Y

*Źródło

orphanet