

DISEASE:
Cerulean cataract

NAME:	Cerulean cataract
DESCRIPTION:	A type of hereditary congenital cataract, distinguished by bluish and white opacifications in the superficial layers of the fetal lens nucleus and adult lens nucleus, and characterized by reduced visual acuity in childhood, eventually necessitating extraction of the lens.
ORPHACODE:	98989
SYNONYMS:	Blue-dot cataract
XREF(S):	Orphanet OMIM OMIM MeSH ICD-10
ANALYTE(S):	CRYBB2 CRYGD MIP MAF
CREATED:	13 May 2019 - 01:02
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