

DISEASE:
Severe early-childhood-onset retinal dystrophy

NAME:	Severe early-childhood-onset retinal dystrophy
DESCRIPTION:	Severe early childhood onset retinal dystrophy (SECORD) is an inherited retinal dystrophy characterized by a severe congenital night blindness, progressive retinal dystrophy and nystagmus. Best corrected visual acuity can reach 0.3 in the first decade of life and can pertain well into the second decade of life. Blindness is often complete by the age of 30 years.
ORPHACODE:	364055
SYNONYMS:	EOSRD Early-onset severe retinal dystrophy SECORD
XREF(S):	<u>Orphanet</u> <u>ICD-10</u>
ANALYTE(S):	<u>RPE65</u> <u>LCA5</u> <u>LRAT</u> <u>SPATA7</u>
CREATED:	13 May 2019 - 01:02
CHANGED:	22 Jun 2023 - 16:14

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