
DISEASE:
Oculocutaneous albinism type 6

NAME:	Oculocutaneous albinism type 6
DESCRIPTION:	A form of oculocutaneous albinism characterized by light hair at birth that darkens with age, white skin, transparent irides, photophobia, nystagmus, foveal hypoplasia and reduced visual acuity.
ORPHACODE:	370097
SYNONYMS:	OCA6
XREF(S):	<u>Orphanet</u> <u>ICD-10</u> <u>OMIM</u>
ANALYTE(S):	<u>SLC24A5</u>
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