

DISEASE:
Retinopathy of prematurity

NAME:	Retinopathy of prematurity
DESCRIPTION:	A rare retinal vasoproliferative disease affecting preterm infants characterized initially by a delay in physiologic retinal vascular development and compromised physiologic vascularity, and subsequently by aberrant angiogenesis in the form of intravitreal neovascularization.
ORPHACODE:	90050
SYNONYMS:	ROP Retrolental fibroplasia
XREF(S):	Orphanet OMIM MedDRA MeSH ICD-10
ANALYTE(S):	LRP5 FZD4 NDP
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