

DISEASE:
Weill-Marchesani syndrome

NAME:	Weill-Marchesani syndrome
DESCRIPTION:	Weill-Marchesani syndrome (WMS) is a rare condition characterized by short stature, brachydactyly, joint stiffness, and characteristic eye abnormalities including microspherophakia, ectopia of the lens, severe myopia, and glaucoma.
ORPHACODE:	3449
SYNONYMS:	Spherophakia-brachymorphia syndrome
XREF(S):	Orphanet MedDRA OMIM OMIM OMIM MeSH ICD-10
ANALYTE(S):	ADAMTS10 FBN1 LTBP2
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