

DISEASE:
Milroy disease

NAME:	Milroy disease
DESCRIPTION:	Milroy disease is a frequent form of primary lymphedema (see this term) characterized generally by painless, chronic lower-limb lymphedema found at birth or developing in the early neonatal period.
ORPHACODE:	79452
SYNONYMS:	Hereditary lymphedema type I Nonne-Milroy lymphedema
XREF(S):	Orphanet ICD-10 OMIM OMIM OMIM OMIM OMIM
ANALYTE(S):	ANGPT2 FLT4 PIEZO1 GJC2
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