
DISEASE:
X-linked severe congenital neutropenia

NAME:	X-linked severe congenital neutropenia
DESCRIPTION:	X-linked severe congenital neutropenia is an immunodeficiency syndrome characterized by recurrent major bacterial infections, severe congenital neutropenia, and monocytopenia. It has been described in five males spanning three generations of one family. It is transmitted as an X-linked recessive trait and is caused by mutations in the WAS gene, encoding the WASP protein.
ORPHACODE:	86788
XREF(S):	Orphanet OMIM ICD-10
ANALYTE(S):	WAS
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