

DISEASE:
Nakajo-Nishimura syndrome

NAME:	Nakajo-Nishimura syndrome
DESCRIPTION:	Nakajo-Nishimura syndrome (NNS) is a rare autoinflammatory disorder belonging to the proteasome disability syndrome (see this term) group, and characterized by pernio-like lesions appearing in infancy followed by recurrent fever, nodular skin eruption, partial lipodystrophy (mainly in upper extremities and face) and joint contractures.
ORPHACODE:	2615
SYNONYMS:	Amyotrophy-fat tissue anomaly syndrome NNS Secondary hypertrophic osteoperiostosis with pernio
XREF(S):	Orphanet ICD-10 OMIM
ANALYTE(S):	PSMB8
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