

DISEASE:
Infantile neuroaxonal dystrophy

NAME:	Infantile neuroaxonal dystrophy
DESCRIPTION:	Infantile neuroaxonal dystrophy/atypical neuroaxonal dystrophy (INAD/atypical NAD) is a type of neurodegeneration with brain iron accumulation (NBIA; see this term) characterized by psychomotor delay and regression, increasing neurological involvement with symmetrical pyramidal tract signs and spastic tetraplegia. INAD may be classic or atypical and patients present with symptoms anywhere along a continuum between the two.
ORPHACODE:	35069
SYNONYMS:	INAD INAD1 PLAN Phospholipase A2-associated neurodegeneration Seitelberger disease
XREF(S):	Orphanet OMIM OMIM ICD-10
ANALYTE(S):	PLA2G6
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