

DISEASE:**Late-onset citrullinemia type I**

NAME:	Late-onset citrullinemia type I
DESCRIPTION:	A form of citrullinemia type I characterized clinically by adult onset of symptoms including variable hyperammonemia and less striking neurological findings which may include intense headache, scotomas, migraine-like episodes, ataxia, slurred speech, lethargy and drowsiness. Serious increased intracranial pressure may occur.
ORPHACODE:	247573
SYNONYMS:	Late-onset citrullinemia type 1
XREF(S):	Orphanet
ANALYTE(S):	ASS1
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