

DISEASE:**Primary hypomagnesemia with secondary hypocalcemia**

NAME:	Primary hypomagnesemia with secondary hypocalcemia
DESCRIPTION:	Primary hypomagnesemia with secondary hypocalcemia (PHSH) is a form of familial primary hypomagnesemia (FPH, see this term), characterized by severe hypomagnesemia and secondary hypocalcemia associated with neurological symptoms, including generalized seizures, tetany and muscle spasms. PHSH may be fatal or may result in chronic irreversible neurological complications.
ORPHACODE:	30924
SYNONYMS:	HOMG1 HSH Hypomagnesemia caused by selective magnesium malabsorption Hypomagnesemia intestinal type 1 Intestinal hypomagnesemia with secondary hypocalcemia PHSH
XREF(S):	Orphanet OMIM ICD-10
ANALYTE(S):	TRPM6
CREATED:	13 May 2019 - 01:02

CHANGED:	22 Jun 2023 - 16:14
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