

DISEASE:**Glycogen storage disease due to glycogen branching enzyme deficiency, progressive hepatic form**

NAME:	Glycogen storage disease due to glycogen branching enzyme deficiency, progressive hepatic form
ORPHACODE:	308621
SYNONYMS:	GBE deficiency, progressive hepatic form GSD due to glycogen branching enzyme deficiency, progressive hepatic form GSD type 4, progressive hepatic form GSDIV, progressive hepatic form Glycogen storage disease type 4, progressive hepatic form Glycogen storage disease type IV, progressive hepatic form Glycogenosis due to glycogen branching enzyme deficiency, progressive hepatic form Glycogenosis type 4, progressive hepatic form Glycogenosis type IV, progressive hepatic form
XREF(S):	Orphanet ICD-10 OMIM
ANALYTE(S):	GBE1
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