

Product datasheet

Recombinant Human GFPT1 protein ab152423

1 Image

Description

Product name	Recombinant Human GFPT1 protein
Expression system	Wheat germ
Accession	Q06210-2
Protein length	Full length protein
Animal free	No
Nature	Recombinant
Species	Human

Sequence

MCGIFAYLNYHVPRTREILETLIKGLQRLEYRGYDSAGVGF
 DGGNDKDW
 EANACKIQLIKKKGKVKALDEEVHKQQDMDLDIEFDVHLGI
 AHTRWATHG
 EPSPVNSHPQRSDDKNEFVIHNGIITNYKDLKKFLESKGYD
 FESETDTE
 TIAKLVKMYDNRESQDTSFTTLVERVIQQLEGAFALVFKS
 VHFPGQAVG
 TRRGSPLLIGVRSEHKLSTDHIPILYRTGKDKKGCNLSRV
 DSTTCLFPV
 EEKAVEYYFASDASAVIEHTNRVIFLEDDDDVAAVVDGRLSI
 HRIKRTAGD
 HPGRAVQTLQMELQQIMKGNFSSFMQKEIFEQPESVVNT
 MRGRVNFDDYT
 VNLGGLKDHKEIQRRCRLILACGTSYHAGVATRQVLEELT
 ELPVMVEL
 ASDFLDRNTPVFRDDVCFFLSQSGETADTLMGLRYCKER
 GALTVGITNTV
 GSSISRETD CGVHINAGPEIGVASTKAYTSQFVSLVMFALM
 MCDDRISMQ
 ERRKEIMLGLKRLPDLIKEVLSMDDEIQKLATELYHQKSVLI
 MGRGYHYA
 TCLEGALKIKEITYMHSEGILAGELKHGPLALVDKLMPVIMII
 MRDHTYA
 KCQNALQQVVARQGRPVICDKEDTETIKNTKRTIKVPHS
 VDCLQGILSV IPLQLLAFHLAVLRGYDVDFPRNLAKSVTVE

Predicted molecular weight 103 kDa including tags

Specifications

Our [Abpromise guarantee](#) covers the use of **ab152423** in the following tested applications.

The application notes include recommended starting dilutions; optimal dilutions/concentrations should be determined by the end user.

Applications	ELISA SDS-PAGE Western blot
Form	Liquid
Additional notes	

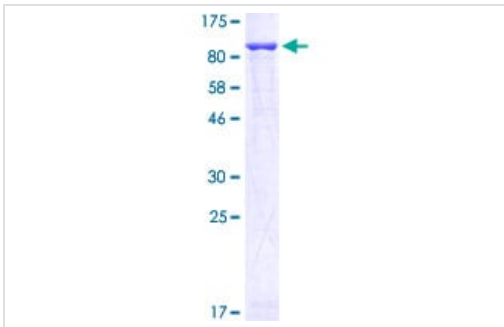
Preparation and Storage

Stability and Storage	Shipped on dry ice. Upon delivery aliquot and store at -80°C. Avoid freeze / thaw cycles. pH: 8.00 Constituents: 0.31% Glutathione, 0.79% Tris HCl
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General Info

Function	Controls the flux of glucose into the hexosamine pathway. Most likely involved in regulating the availability of precursors for N- and O-linked glycosylation of proteins.
Tissue specificity	Isoform 1 is predominantly expressed in skeletal muscle. Not expressed in brain. Seems to be selectively expressed in striated muscle.
Pathway	Nucleotide-sugar biosynthesis; UDP-N-acetyl-alpha-D-glucosamine biosynthesis; alpha-D-glucosamine 6-phosphate from D-fructose 6-phosphate: step 1/1.
Involvement in disease	Defects in GFPT1 are the cause of limb-girdle myasthenia with tubular aggregates (LGMTA) [MIM:610542]. A congenital myasthenic syndrome characterized by onset of proximal muscle weakness in the first decade. Individuals with this condition have a recognizable pattern of weakness of shoulder and pelvic girdle muscles, and sparing of ocular or facial muscles. EMG classically shows a decremental response to repeated nerve stimulation, a sign of neuromuscular junction dysfunction. Affected individuals show a favorable response to acetylcholinesterase (AChE) inhibitors.
Sequence similarities	Contains 1 glutamine amidotransferase type-2 domain. Contains 2 SIS domains.

Images



12.5% SDS-PAGE analysis of ab152423 stained with Coomassie Blue.

SDS-PAGE - Recombinant Human GFPT1 protein
(ab152423)

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