

Product datasheet for TP301304

OriGene Technologies, Inc.

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Galactosidase alpha (GLA) (NM_000169) Human Recombinant Protein

Product data:

Product Type: Recombinant Proteins

Description: Recombinant protein of human galactosidase, alpha (GLA), 20 μg

Species: Human Expression Host: HEK293T

Expression cDNA >RC201304 protein sequence
Clone or AA Sequence: Red=Cloning site Green=Tags(s)

MQLRNPELHLGCALALRFLALVSWDIPGARALDNGLARTPTMGWLHWERFMCNLDCQEEPDSCISEKLFM EMAELMVSEGWKDAGYEYLCIDDCWMAPQRDSEGRLQADPQRFPHGIRQLANYVHSKGLKLGIYADVGNK TCAGFPGSFGYYDIDAQTFADWGVDLLKFDGCYCDSLENLADGYKHMSLALNRTGRSIVYSCEWPLYMWP FQKPNYTEIRQYCNHWRNFADIDDSWKSIKSILDWTSFNQERIVDVAGPGGWNDPDMLVIGNFGLSWNQQ VTQMALWAIMAAPLFMSNDLRHISPQAKALLQDKDVIAINQDPLGKQGYQLRQGDNFEVWERPLSGLAWA VAMINRQEIGGPRSYTIAVASLGKGVACNPACFITQLLPVKRKLGFYEWTSRLRSHINPTGTVLLQLENT

MQMSLKDLL

TRTRPLEQKLISEEDLAANDILDYKDDDDKV

Tag: C-Myc/DDK
Predicted MW: 45.3 kDa

Concentration: $>0.05 \mu g/\mu L$ as determined by microplate BCA method

Purity: > 80% as determined by SDS-PAGE and Coomassie blue staining

Buffer: 25 mM Tris-HCl, 100 mM glycine, pH 7.3, 10% glycerol

Preparation: Recombinant protein was captured through anti-DDK affinity column followed by conventional

chromatography steps.

Note: For testing in cell culture applications, please filter before use. Note that you may experience

some loss of protein during the filtration process.

Storage: Store at -80°C.

Stability: Stable for 12 months from the date of receipt of the product under proper storage and handling

conditions. Avoid repeated freeze-thaw cycles.

RefSeq: NP 000160





Locus ID: 2717

UniProt ID: <u>P06280</u>, <u>Q53Y83</u>

RefSeq Size: 1418
Cytogenetics: Xq22.1
RefSeq ORF: 1288
Synonyms: GALA

Summary: This gene encodes a homodimeric glycoprotein that hydrolyses the terminal alpha-galactosyl

moieties from glycolipids and glycoproteins. This enzyme predominantly hydrolyzes ceramide trihexoside, and it can catalyze the hydrolysis of melibiose into galactose and glucose. A variety of mutations in this gene affect the synthesis, processing, and stability of this enzyme, which causes Fabry disease, a rare lysosomal storage disorder that results from a failure to catabolize

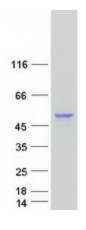
alpha-D-galactosyl glycolipid moieties. [provided by RefSeq, Jul 2008]

Protein Families: Druggable Genome

Protein Pathways: Galactose metabolism, Glycerolipid metabolism, Glycosphingolipid biosynthesis - globo series,

Lysosome, Sphingolipid metabolism

Product images:



Coomassie blue staining of purified GLA protein (Cat# TP301304). The protein was produced from HEK293T cells transfected with GLA cDNA clone (Cat# [RC201304]) using MegaTran 2.0 (Cat# [TT210002]).