

Product datasheet for TP301614L

GBA (NM_001005741) Human Recombinant Protein

Product data:

OriGene Technologies, Inc.

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Product Type:	Recombinant Proteins	
Description:	Purified recombinant protein of Homo sapiens glucosidase, beta; acid (includes glucosylceramidase) (GBA), transcript variant 2, 1 mg	
Species:	Human	
Expression Host:	HEK293T	
Expression cDNA Clone or AA Sequence:	>RC201614 protein sequence Red=Cloning site Green=Tags(s)	
	MEFSSPSREECPKPLSRVSIMAGSLTGLLLLQAVSWASGARPCIPKSFGYSSVVCVCNATYCDSFDPPTF PALGTFSRYESTRSGRRMELSMGPIQANHTGTGLLLTLQPEQKFQKVKGFGGAMTDAAALNILALSPPAQ NLLLKSYFSEEGIGYNIIRVPMASCDFSIRTYTYADTPDDFQLHNFSLPEEDTKLKIPLIHRALQLAQRP VSLLASPWTSPTWLKTNGAVNGKGSLKGQPGDIYHQTWARYFVKFLDAYAEHKLQFWAVTAENEPSAGLL SGYPFQCLGFTPEHQRDFIARDLGPTLANSTHHNVRLLMLDDQRLLLPHWAKVVLTDPEAAKYVHGIAVH WYLDFLAPAKATLGETHRLFPNTMLFASEACVGSKFWEQSVRLGSWDRGMQYSHSIITNLLYHVVGWTDW NLALNPEGGPNWVRNFVDSPIIVDITKDTFYKQPMFYHLGHFSKFIPEGSQRVGLVASQKNDLDAVALMH PDGSAVVVVLNRSSKDVPLTIKDPAVGFLETISPGYSIHTYLWRRQ	
	TRTRPLEQKLISEEDLAANDILDYKDDDDKV	
Tag:	C-Myc/DDK	
Predicted MW:	55.5 kDa	
Concentration:	>0.05 µg/µL 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.	



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	GBA (NM_001005741) Human Recombinant Protein – TP301614L Stable for 12 months from the date of receipt of the product under proper storage and handling conditions. Avoid repeated freeze-thaw cycles.	
Stability:		
RefSeq:	<u>NP 001005741</u>	
Locus ID:	2629	
UniProt ID:	<u>P04062, B7Z6S9, A0A068F658</u>	
RefSeq Size:	2583	
Cytogenetics:	1q22	
RefSeq ORF:	1608	
Synonyms:	GBA1; GCB; GLUC	
Summary:	This gene encodes a lysosomal membrane protein that cleaves the beta-glucosidic linkage of glycosylceramide, an intermediate in glycolipid metabolism. Mutations in this gene cause Gaucher disease, a lysosomal storage disease characterized by an accumulation of glucocerebrosides. A related pseudogene is approximately 12 kb downstream of this gene on chromosome 1. Alternative splicing results in multiple transcript variants. [provided by RefSeq, Jan 2010]	
Protein Families	: Druggable Genome	
Protein Pathway	Lysosome, Metabolic pathways, Other glycan degradation, Sphingolipid metabolism	

Product images:

116 -	-	
66 -	-	-
45 -	-	-
35 -	-	
25 -	-	
18 -	-	
14 -	-1	

Coomassie blue staining of purified GBA protein (Cat# [TP301614]). The protein was produced from HEK293T cells transfected with GBA cDNA clone (Cat# [RC201614]) using MegaTran 2.0 (Cat# [TT210002]).

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