

Product datasheet for TP327162L

SPART (NM_001142295) Human Recombinant Protein

Product data:

Storage:

Product Type: Recombinant Proteins Description: Purified recombinant protein of Homo sapiens spastic paraplegia 20 (Troyer syndrome) (SPG20), transcript variant 3, 1 mg Species: Human **Expression Host:** HEK293T Expression cDNA Clone >RC227162 protein sequence Red=Cloning site Green=Tags(s) or AA Sequence: MEQEPQNGEPAEIKIIREAYKKAFLFVNKGLNTDELGQKEEAKNYYKQGIGHLLRGISISSKESEHTGTG WESARQMQQKMKETLQNVRTRLEILEKGLATSLQNDLQEVPKLYPEFPPKDMCEKLPEPQSFSSAPQHAE VNGNTSTPSAGAVAAPASLSLPSQSCPAEAPPAYTPQAAEGHYTVSYGTDSGEFSSVGEEFYRNHSQPPP LETLGLDADELILIPNGVQIFFVNPAGEVSAPSYPGYLRIVRFLDNSLDTVLNRPPGFLQVCDWLYPLVP DRSPVLKCTAGAYMFPDTMLQAAGCFVGVVLSSELPEDDRELFEDLLRQMSDLRLQANWNRAEEENEFQI PGRTRPSSDQLKEASGTDVKQLDQGNKDVRHKGKRGKRAKDTSSEEVNLSHIVPCEPVPEEKPKELHEWS EKVAHNILSGASWVSWGLVKGAEITGKAIQKGASKLRERIQPEEKPVEVSPAVTKGLYIAKQATGGAAKV SQFLVDGVCTVANCVGKELAPHVKKHGSKLVPESLKKDKDGKSPLDGAMVVAASSVQGFSTVWQGLECAA KCIVNNVSAETVQTVRYKYGYNAGEATHHAVDSAVNVGVTAYNINNIGIKAMVKKTATQTGHTLLEDYQI VDNSQRENQEGAANVNVRGEKDEQTKEVKEAKKKDK **TRTRPLEQKLISEEDLAANDILDYKDDDDKV** Tag: C-Myc/DDK Predicted MW: 72.7 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. For testing in cell culture applications, please filter before use. Note that you may experience Note: some loss of protein during the filtration process.



This product is to be used for laboratory only. Not for diagnostic or therapeutic use. ©2023 OriGene Technologies, Inc., 9620 Medical Center Drive, Ste 200, Rockville, MD 20850, US

Store at -80°C.

OriGene Technologies, Inc.

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	SPART (NM_001142295) Human Recombinant Protein – TP327162L	
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:	<u>NP 001135767</u>	
Locus ID:	23111	
UniProt ID:	<u>Q8N0X7</u> , <u>A0A024RDV9</u>	
RefSeq Size:	4962	
Cytogenetics:	13q13.3	
RefSeq ORF:	1998	
Synonyms:	SPG20; TAHCCP1	
Summary:	This gene encodes a protein containing a MIT (Microtubule Interacting and Trafficking molecule) domain, and is implicated in regulating endosomal trafficking and mitochondria function. The protein localizes to mitochondria and partially co-localizes with microtubules. Stimulation with epidermal growth factor (EGF) results in protein translocation to the plasma membrane, and the protein functions in the degradation and intracellular trafficking of EGF receptor. Multiple alternatively spliced variants, encoding the same protein, have been identified. Mutations associated with this gene cause autosomal recessive spastic paraplegia 20 (Troyer syndrome). [provided by RefSeq, Nov 2008]	

Product images:

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

Coomassie blue staining of purified SPART protein (Cat# [TP327162]). The protein was produced from HEK293T cells transfected with SPART cDNA clone (Cat# [RC227162]) using MegaTran 2.0 (Cat# [TT210002]).

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