

# Product datasheet for TP325431L

### SPG21 (NM\_001127889) Human Recombinant Protein

### **Product data:**

#### **Product Type: Recombinant Proteins Description:** Recombinant protein of human spastic paraplegia 21 (autosomal recessive, Mast syndrome) (SPG21), transcript variant 2, 1 mg Species: Human **Expression Host:** HEK293T Expression cDNA Clone >RC225431 protein sequence or AA Sequence: Red=Cloning site Green=Tags(s) MGEIKVSPDYNWFRGTVPLKKIIVDDDDSKIWSLYDAGPRSIRCPLIFLPPVSGTADVFFRQILALTGWG YRVIALQYPVYWDHLEFCDGFRKLLDHLQLDKVHLFGASLGGFLAQKFAEYTHKSPRVHSLILCNSFSDT SIFNQTWTANSFWLMPAFMLKKIVLGNFSSGPVDPMMADAIDFMVDRLESLGQSELASRLTLNCQNSYVE PHKIRDIPVTIMDVFDQSALSTEAKEEMYKLYPNARRAHLKTGGNFPYLCRSAEVNLYVQIHLLQFHGTK YAAIDPSMVSAEELEVQKGSLGISQEEQ **TRTRPL**EQKLISEEDLAANDILDYKDDDDKV Tag: C-Myc/DDK Predicted MW: 34.8 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. For testing in cell culture applications, please filter before use. Note that you may experience Note: 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 001121361 Locus ID: 51324



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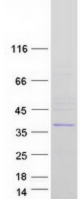
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### OriGene Technologies, Inc.

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	SPG21 (NM_001127889) Human Recombinant Protein – TP325431L
UniProt ID:	<u>Q9NZD8, A0A024R5Y1</u>
RefSeq Size:	1658
Cytogenetics:	15q22.31
RefSeq ORF:	924
Synonyms:	ABHD21; ACP33; BM-019; GL010; MAST
Summary:	The protein encoded by this gene binds to the hydrophobic C-terminal amino acids of CD4 which are involved in repression of T cell activation. The interaction with CD4 is mediated by the noncatalytic alpha/beta hydrolase fold domain of this protein. It is thus proposed that this gene product modulates the stimulatory activity of CD4. Mutations in this gene are associated with autosomal recessive spastic paraplegia 21 (SPG21), also known as mast syndrome. Alternative splicing results in multiple transcript variants. [provided by RefSeq, Aug 2014]

## **Product images:**



Coomassie blue staining of purified SPG21 protein (Cat# [TP325431]). The protein was produced from HEK293T cells transfected with SPG21 cDNA clone (Cat# [RC225431]) using MegaTran 2.0 (Cat# [TT210002]).

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