

Product datasheet for TP319188

OriGene Technologies, Inc.

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VPS13B (NM_181661) Human Recombinant Protein

Product data:

Product Type: Recombinant Proteins

Description: Recombinant protein of human vacuolar protein sorting 13 homolog B (yeast) (VPS13B),

transcript variant 4, 20 µg

Species: Human Expression Host: HEK293T

Expression cDNA Clone >RC219188 representing NM_181661 **or AA Sequence:** Red=Cloning site Green=Tags(s)

MLESYVTPILMSYVNRYIKNLKPSDLQLSLWGGDVVLSKLELKLDVLEQELKLPFTFLSGHIHELRIHVP WTKLGSEPVVITINTMECILKLKDGIQDDHESCGSNSTNRSTAESTKSSIKPRRMQQAAPTDPDLPPGYV QSLIRRVVNNVNIVINNLILKYVEDDIVLSVNITSAECYTVGELWDRAFMDISATDLVLRKVINFSDCTV CLDKRNASGKIEFYQDPLLYKCSFRTRLHFTYENLNSKMPSVIKIHTLVESLKLSITDQQLPMFIRIMQL GIALYYGEIGNFKEGEIEDLTCHNKDMLGNITGSEDETRIDMQYPAQHKGQELYSQQDEEQPQGWVSWAW

SFVPAIVSYDDGEEDFVGNDPASTMHQQKAQTLKDPIVSIGFYCTKATVTFKVGLFSCCLYLYQL

TRTRPLEQKLISEEDLAANDILDYKDDDDKV

Tag: C-Myc/DDK

Predicted MW: 47 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.

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 858047





RefSeq ORF:

Locus ID: 157680

UniProt ID: Q7Z7G8 RefSeq Size: 1634 Cytogenetics: 8q22.2

Synonyms: CHS1; COH1

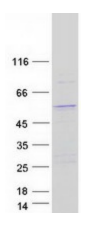
1245

Summary: This gene encodes a potential transmembrane protein that may function in vesicle-mediated

transport and sorting of proteins within the cell. This protein may play a role in the

development and the function of the eye, hematological system, and central nervous system. Mutations in this gene have been associated with Cohen syndrome. Multiple splice variants encoding distinct isoforms have been identified for this gene. [provided by RefSeq, Jul 2008]

Product images:



Coomassie blue staining of purified VPS13B protein (Cat# TP319188). The protein was produced from HEK293T cells transfected with VPS13B cDNA clone (Cat# [RC219188]) using

MegaTran 2.0 (Cat# [TT210002]).