

Product datasheet for TP303457M

OriGene Technologies, Inc.

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WASP (WAS) (NM_000377) Human Recombinant Protein

Product data:

Product Type: Recombinant Proteins

Description: Recombinant protein of human Wiskott-Aldrich syndrome (eczema-thrombocytopenia) (WAS),

100 µg

Species: Human Expression Host: HEK293T

Expression cDNA >RC203457 representing NM_000377

Clone or AA Sequence: Red=Cloning site Green=Tags(s)

MSGGPMGGRPGGRGAPAVQQNIPSTLLQDHENQRLFEMLGRKCLTLATAVVQLYLALPPGAEHWTKEHCG AVCFVKDNPQKSYFIRLYGLQAGRLLWEQELYSQLVYSTPTPFFHTFAGDDCQAGLNFADEDEAQAFRAL VQEKIQKRNQRQSGDRRQLPPPPTPANEERRGGLPPLPLHPGGDQGGPPVGPLSLGLATVDIQNPDITSS RYRGLPAPGPSPADKKRSGKKKISKADIGAPSGFKHVSHVGWDPQNGFDVNNLDPDLRSLFSRAGISEAQ LTDAETSKLIYDFIEDQGGLEAVRQEMRRQEPLPPPPPSRGGNQLPRPPIVGGNKGRSGPLPPVPLGIA PPPPTPRGPPPPGRGGPPPPPPATGRSGPLPPPPPGAGGPPMPPPPPPPPSSGNGPAPPPLPPALV PAGGLAPGGGRGALLDQIRQGIQLNKTPGAPESSALQPPPQSSEGLVGALMHVMQKRSRAIHSSDEGEDQ

AGDEDEDDEWDD

TRTRPLEQKLISEEDLAANDILDYKDDDDKV

Tag: C-Myc/DDK

Predicted MW: 52.7 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.

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.





WASP (WAS) (NM_000377) Human Recombinant Protein - TP303457M

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 000368

Locus ID: 7454

UniProt ID: <u>P42768</u>, <u>A0A024QYX8</u>

RefSeq Size: 1806 Cytogenetics: Xp11.23 RefSeq ORF: 1506

Synonyms: IMD2; SCNX; THC; THC1; WASP; WASPA

Summary: The Wiskott-Aldrich syndrome (WAS) family of proteins share similar domain structure, and are

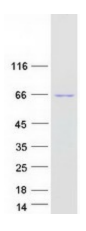
involved in transduction of signals from receptors on the cell surface to the actin cytoskeleton. The presence of a number of different motifs suggests that they are regulated by a number of different stimuli, and interact with multiple proteins. Recent studies have demonstrated that these proteins, directly or indirectly, associate with the small GTPase, Cdc42, known to regulate formation of actin filaments, and the cytoskeletal organizing complex, Arp2/3. Wiskott-Aldrich syndrome is a rare, inherited, X-linked, recessive disease characterized by immune dysregulation and microthrombocytopenia, and is caused by mutations in the WAS gene. The WAS gene product is a cytoplasmic protein, expressed exclusively in hematopoietic cells, which show signalling and cytoskeletal abnormalities in WAS patients. A transcript variant arising as a result of alternative promoter usage, and containing a different 5' UTR sequence, has been described, however, its full-length nature is not known. [provided by RefSeq, Jul 2008]

Protein Families: Druggable Genome

Protein Pathways: Adherens junction, Chemokine signaling pathway, Fc gamma R-mediated phagocytosis,

Pathogenic Escherichia coli infection, Regulation of actin cytoskeleton

Product images:



Coomassie blue staining of purified WAS protein (Cat# [TP303457]). The protein was produced from HEK293T cells transfected with WAS cDNA clone (Cat# [RC203457]) using MegaTran 2.0 (Cat# [TT210002]).