

# Product datasheet for TP304771L

## TIMM8A (NM\_004085) Human Recombinant Protein

### **Product data:**

#### **Product Type: Recombinant Proteins** Recombinant protein of human translocase of inner mitochondrial membrane 8 homolog A **Description:** (yeast) (TIMM8A), nuclear gene encoding mitochondrial protein, transcript variant 1, 1 mg Species: Human **Expression Host:** HEK293T **Expression cDNA Clone** >RC204771 protein sequence or AA Sequence: Red=Cloning site Green=Tags(s) MDSSSSSSAAGLGAVDPQLQHFIEVETQKQRFQQLVHQMTELCWEKCMDKPGPKLDSRAEACFVNCVE RF IDTSQFILNRLEQTQKSKPVFSESLSD **TRTRPLEQKLISEEDLAANDILDYKDDDDKV** Tag: C-Myc/DDK Predicted MW: 10.8 kDa Concentration: >0.05 µg/µL as determined by microplate BCA method > 80% as determined by SDS-PAGE and Coomassie blue staining **Purity: Buffer:** 25 mM Tris-HCl, 100 mM glycine, pH 7.3, 10% glycerol Recombinant protein was captured through anti-DDK affinity column followed by **Preparation:** 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. Store at -80°C. Storage: Stable for 12 months from the date of receipt of the product under proper storage and Stability: handling conditions. Avoid repeated freeze-thaw cycles. RefSeq: NP 004076 1678 Locus ID: **UniProt ID:** 060220



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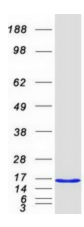
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	TIMM8A (NM_004085) Human Recombinant Protein – TP304771L
RefSeq Size:	1459
Cytogenetics:	Xq22.1
RefSeq ORF:	291
Synonyms:	DDP; DDP1; DFN1; MTS; TIM8
Summary:	This translocase is involved in the import and insertion of hydrophobic membrane proteins from the cytoplasm into the mitochondrial inner membrane. The gene is mutated in Mohr- Tranebjaerg syndrome/Deafness Dystonia Syndrome (MTS/DDS) and it is postulated that MTS/DDS is a mitochondrial disease caused by a defective mitochondrial protein import system. Defects in this gene also cause Jensen syndrome; an X-linked disease with opticoacoustic nerve atrophy and muscle weakness. This protein, along with TIMM13, forms a 70 kDa heterohexamer. Alternative splicing results in multiple transcript variants encoding distinct isoforms.[provided by RefSeq, Mar 2009]
Protein Families	: Druggable Genome

Product images:



Coomassie blue staining of purified TIMM8A protein (Cat# [TP304771]). The protein was produced from HEK293T cells transfected with TIMM8A cDNA clone (Cat# [RC204771]) using MegaTran 2.0 (Cat# [TT210002]).

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