

# **Product datasheet for TP319648M**

#### OriGene Technologies, Inc.

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### Tropomyosin 2 (TPM2) (NM\_003289) Human Recombinant Protein

**Product data:** 

**Product Type:** Recombinant Proteins

**Description:** Recombinant protein of human tropomyosin 2 (beta) (TPM2), transcript variant 1, 100 μg

Species: Human
Expression Host: HEK293T

**Expression cDNA Clone** >RC219648 representing NM\_003289 or AA Sequence: Red=Cloning site Green=Tags(s)

MDAIKKKMQMLKLDKENAIDRAEQAEADKKQAEDRCKQLEEEQQALQKKLKGTEDEVEKYSESVKEAQEK LEQAEKKATDAEADVASLNRRIQLVEEELDRAQERLATALQKLEEAEKAADESERGMKVIENRAMKDEEK MELQEMQLKEAKHIAEDSDRKYEEVARKLVILEGELERSEERAEVAESKCGDLEEELKIVTNNLKSLEAQ ADKYSTKEDKYEEEIKLLEEKLKEAETRAEFAERSVAKLEKTIDDLEDEVYAQKMKYKAISEELDNALND

ITSL

**TRTRPLEQKLISEEDLAANDILDYKDDDDKV** 

Tag: C-Myc/DDK
Predicted MW: 32.7 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 003280

**Locus ID:** 7169





#### Tropomyosin 2 (TPM2) (NM\_003289) Human Recombinant Protein - TP319648M

**UniProt ID:** P07951 RefSeq Size: 1327 Cytogenetics: 9p13.3 RefSeq ORF: 852

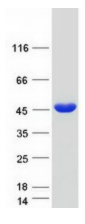
Synonyms: AMCD1; DA1; DA2B; DA2B4; HEL-S-273; NEM4; TMSB

**Summary:** This gene encodes beta-tropomyosin, a member of the actin filament binding protein family,

> and mainly expressed in slow, type 1 muscle fibers. Mutations in this gene can alter the expression of other sarcomeric tropomyosin proteins, and cause cap disease, nemaline myopathy and distal arthrogryposis syndromes. Alternatively spliced transcript variants encoding different isoforms have been found for this gene.[provided by RefSeq, Mar 2009]

Cardiac muscle contraction, Dilated cardiomyopathy, Hypertrophic cardiomyopathy (HCM) **Protein Pathways:** 

## **Product images:**



Coomassie blue staining of purified TPM2 protein (Cat# [TP319648]). The protein was produced from HEK293T cells transfected with TPM2 cDNA clone (Cat# [RC219648]) using MegaTran 2.0

(Cat# [TT210002]).