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Lieferung & Zahlungsart

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Zuschläge

- Mindermengenzuschlag
- Trockeneiszuschlag
- Gefahrgutzuschlag
- Expressversand

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DTNA 293T Cell Transient Overexpression Lysate(Denatured)

Catalog # : H00001837-T01

規格 : [100 uL]

[List All](#)

Specification

Transfected Cell Line: 293T

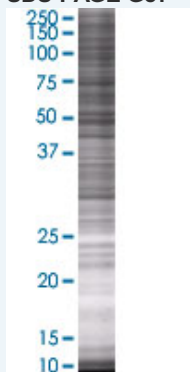
Plasmid: pCMV-DTNA full-length

Host: Human

Theoretical MW (kDa): 75.57

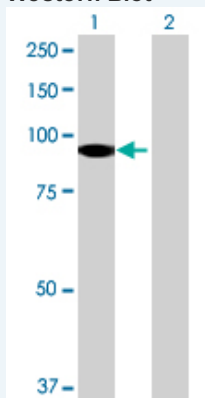
Quality Control Testing: Transient overexpression cell lysate was tested with Anti-DTNA antibody (H00001837-B02) by Western Blots.

SDS-PAGE Gel



DTNA transfected lysate.

Western Blot



Lane 1: DTNA transfected lysate (75.57 KDa)

Lane 2: Non-transfected lysate.

Storage Buffer: 1X Sample Buffer (50 mM Tris-HCl, 2% SDS, 10% glycerol, 300 mM 2-mercaptoethanol, 0.01% Bromophenol blue)

Storage Instruction: Store at -80°C. Aliquot to avoid repeated freezing and thawing.

MSDS:  [Download](#)

Applications

Western Blot

Gene Information

Entrez GeneID: [1837](#)

GeneBank
Accession#: [NM_032975.2](#)

Protein
Accession#: [NP_116757.2](#)

Gene Name: DTNA

Gene Alias: D18S892E,DRP3,DTN,FLJ96209,LVNC1

Gene
Description: dystrobrevin, alpha

Omim ID: [601239](#), [604169](#), [606617](#)

Gene Ontology: [Hyperlink](#)

Gene Summary: The protein encoded by this gene belongs to the dystrobrevin subfamily of the dystrophin family. This protein is a component of the dystrophin-associated protein complex (DPC), which consists of dystrophin and several integral and peripheral membrane proteins, including dystroglycans, sarcoglycans, syntrophins and alpha- and beta-dystrobrevin. The DPC localizes to the sarcolemma and its disruption is associated with various forms of muscular dystrophy. Mutations in this gene are associated with left ventricular noncompaction with congenital heart defects. Multiple alternatively spliced transcript variants encoding different isoforms have been identified for this gene. [provided by RefSeq]

Other
Designations: OTTHUMP00000163154,OTTHUMP00000163155,dystrobrevin alpha,dystrophin-related protein 3

Related Disease

[Tobacco Use Disorder](#)

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