

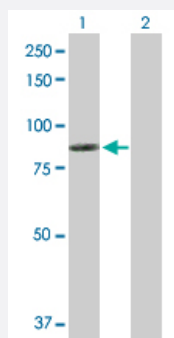
MaxPab®

DMD purified MaxPab rabbit polyclonal antibody (D01P)

Catalog # H00001756-D01P

Size 100 ug

Applications



Western Blot (Transfected lysate)

Western Blot analysis of DMD expression in transfected 293T cell line ([H00001756-T01](#)) by DMD MaxPab polyclonal antibody.

Lane 1: DMD transfected lysate(72.20 KDa).

Lane 2: Non-transfected lysate.

Specification

Product Description

Rabbit polyclonal antibody raised against a full-length human DMD protein.

Immunogen

DMD (NP_004007.1, 1 a.a. ~ 635 a.a) full-length human protein.

Sequence

MREQLKGHETQTTTCWDHPKMTELYQSLADLNNVRFSA YRTAMKLRRQLKALCLDLLSLAACDA
LDQHNLKQNDQPM DILQIINCLTTIYDRLEQEHNNLVNVLPCVDMCLNWLLNVYDTGRTGRIRVLSF
KTGIISLCKAHLEDKYRYLFKQVASSTGFCDQRRGLLLHDSIQIPRQLGEVASFGGSNIEPSVRSC
FQFANNKPEIEAALFLDWMRLEPQSMVWLPVLHRVAAAETAKHQAKCNICKECP IIGFRYRSLKH
FNYDICQSCFFSGRVAKGHKMHYPMVEYCTPTTSGEDVRDFAKVLKNKFRTKRYFAKHPRMGYL
PVQTVLEGDNMETPVT LINFWPVDSAPASSPQLSHDDTHSRIEHYASRLAEMENSNGSYLND SIS
PNESIDDEHLLIQHYCQSLNQDSPLSQPRSPAQILISLESEERGELEERILADLEENRNLQAEYDRL
KQQHEHKGLSPLPSPPEMMPTSPQSPRDAELIAEAKLLRQHKGRLEARMQILEDH NKQLESQ L H
RLRQLLEQPQAEAKVNGTTVSSPSTSLQRSDSSQPM LLRVVGSQTSDSMGEEDLLSPPQDTST
GLEEVMEQLNNSFPSSRGHNVGSLFHMADDLGRAMESLVSVMTDEEGAE

Host

Rabbit

Reactivity

Human

Interspecies Antigen Sequence

Mouse (99)

Quality Control Testing	Antibody reactive against mammalian transfected lysate.
Storage Buffer	In 1x PBS, pH 7.4
Storage Instruction	Store at -20°C or lower. Aliquot to avoid repeated freezing and thawing.

Applications

- Western Blot (Transfected lysate)

Western Blot analysis of DMD expression in transfected 293T cell line ([H00001756-T01](#)) by DMD MaxPab polyclonal antibody.

Lane 1: DMD transfected lysate(72.20 KDa).

Lane 2: Non-transfected lysate.

[Protocol Download](#)

Gene Info — DMD

Entrez GeneID	1756
GeneBank Accession#	NM_004016.1
Protein Accession#	NP_004007.1
Gene Name	DMD
Gene Alias	BMD, CMD3B, DXS142, DXS164, DXS206, DXS230, DXS239, DXS268, DXS269, DXS270, DXS272
Gene Description	dystrophin
Omim ID	300376 300377 302045 310200
Gene Ontology	Hyperlink

Gene Summary

The dystrophin gene is the largest gene found in nature, measuring 2.4 Mb. The gene was identified through a positional cloning approach, targeted at the isolation of the gene responsible for Duchenne (DMD) and Becker (BMD) Muscular Dystrophies. DMD is a recessive, fatal, X-linked disorder occurring at a frequency of about 1 in 3,500 new-born males. BMD is a milder allelic form. In general, DMD patients carry mutations which cause premature translation termination (nonsense or frame shift mutations), while in BMD patients dystrophin is reduced either in molecular weight (derived from in-frame deletions) or in expression level. The dystrophin gene is highly complex, containing at least eight independent, tissue-specific promoters and two polyA-addition sites. Furthermore, dystrophin RNA is differentially spliced, producing a range of different transcripts, encoding a large set of protein isoforms. Dystrophin (as encoded by the Dp427 transcripts) is a large, rod-like cytoskeletal protein which is found at the inner surface of muscle fibers. Dystrophin is part of the dystrophin-glycoprotein complex (DGC), which bridges the inner cytoskeleton (F-actin) and the extra-cellular matrix. [provided by RefSeq]

Other Designations

Duchenne muscular dystrophy protein|OTTHUMP00000023117|OTTHUMP00000023124|OTTHUMP00000023125|OTTHUMP00000023126|muscular dystrophy, Duchenne and Becker types

Pathway

- [Arrhythmogenic right ventricular cardiomyopathy \(ARVC\)](#)
- [Hypertrophic cardiomyopathy \(HCM\)](#)

Disease

- [Bone Marrow Diseases](#)
- [Cardiomyopathy](#)
- [Cardiovascular Diseases](#)
- [Cognition](#)
- [Cognition Disorders](#)
- [Diabetes Mellitus](#)
- [Duchenne muscular dystrophy](#)
- [Edema](#)
- [Genetic Predisposition to Disease](#)
- [Intelligence Tests](#)
- [Mental Retardation](#)

- [Muscular Dystrophies](#)
- [Muscular Dystrophy](#)
- [Neuropsychological Tests](#)
- [Ventricular Dysfunction](#)