

Resource Summary Report

Generated by [RRID](#) on Aug 7, 2026

Mouse Anti-Human Dystrophin Monoclonal Antibody, Unconjugated Conjugated, Clone Dy8/6C5

RRID:AB_442081

Type: Antibody

Proper Citation

Leica Biosystems Cat# NCL-DYS2, RRID:AB_442081

Antibody Information

URL: http://antibodyregistry.org/AB_442081

Proper Citation: Leica Biosystems Cat# NCL-DYS2, RRID:AB_442081

Target Antigen: Human Dystrophin

Host Organism: mouse

Clonality: monoclonal

Comments: manufacturer recommendations: Immunohistochemistry; Western Blot; IHC (Frozen), Western Blot

Antibody Name: Mouse Anti-Human Dystrophin Monoclonal Antibody, Unconjugated Conjugated, Clone Dy8/6C5

Description: This monoclonal targets Human Dystrophin

Target Organism: rat, hamster, canine, mouse, rabbit, human

Clone ID: Clone Dy8/6C5

Antibody ID: AB_442081

Vendor: Leica Biosystems

Catalog Number: NCL-DYS2

Record Creation Time: 20250820T040226+0000

Record Last Update: 20250820T175214+0000

Ratings and Alerts

No rating or validation information has been found for Mouse Anti-Human Dystrophin Monoclonal Antibody, Unconjugated Conjugated, Clone Dy8/6C5.

No alerts have been found for Mouse Anti-Human Dystrophin Monoclonal Antibody, Unconjugated Conjugated, Clone Dy8/6C5.

Data and Source Information

Source: [Antibody Registry](#)

Usage and Citation Metrics

We found 9 mentions in open access literature.

Listed below are recent publications. The full list is available at [RRID](#) .

Mozin E, et al. (2024) Dystrophin deficiency impairs cell junction formation during embryonic myogenesis from pluripotent stem cells. *iScience*, 27(7), 110242.

Gorokhova S, et al. (2023) Unusually severe muscular dystrophy upon in-frame deletion of the dystrophin rod domain and lack of compensation by membrane-localized utrophin. *Med (New York, N.Y.)*, 4(4), 245.

Mozin E, et al. (2023) Dystrophin deficiency impairs cell junction formation during embryonic myogenesis. *bioRxiv : the preprint server for biology*.

Noviello C, et al. (2022) RhoA within myofibers controls satellite cell microenvironment to allow hypertrophic growth. *iScience*, 25(1), 103616.

McKee CM, et al. (2022) The anti-aging protein Klotho affects early postnatal myogenesis by downregulating Jmjd3 and the canonical Wnt pathway. *FASEB journal : official publication of the Federation of American Societies for Experimental Biology*, 36(3), e22192.

Taglietti V, et al. (2022) Duchenne muscular dystrophy trajectory in R-DMDdel52 preclinical rat model identifies COMP as biomarker of fibrosis. *Acta neuropathologica communications*, 10(1), 60.

Steinert ND, et al. (2021) Mapping of the contraction-induced phosphoproteome identifies TRIM28 as a significant regulator of skeletal muscle size and function. *Cell reports*, 34(9), 108796.

Hunter DD, et al. (2019) CNS synapses are stabilized trans-synaptically by laminins and laminin-interacting proteins. *The Journal of comparative neurology*, 527(1), 67.

Debashree B, et al. (2018) Mitochondrial dysfunction in human skeletal muscle biopsies of lipid storage disorder. *Journal of neurochemistry*, 145(4), 323.