

Product Data Sheet: mCherry Mouse Monoclonal Antibody

Cat. No: MAB-94019
Conjugate: Unconjugated

Size: 100 ug
Clone: 1C51
Concentration: 1mg/ml
Host: Mouse

Isotype: IgG2a heavy, κ light

Immunogen: Full length recombinant protein

Reactivity: All Species

Western Blot: 1:2,000

Applications: Immunofluorescence: 1:500

Immunohisotchemistry: 1:500

Molecular Weight: ~28kDa **Purification:** Purified

mCherry protein is derived from a natural product, DsRed, originally isolated as a red fluorescent protein from the coral of the genus Discosoma (1). As with other

natural fluorescent proteins of

Cnidarians (jelly fish, sea anemones, and corals), the natural form of the protein forms stable tetramers in vivo. DsRed was engineered to improve its spectral properties and also prevent ultimerization in the Tsien lab, where much work on fluorescent proteins was performed (2). Roger Tsien, along with Martin Chalfie,

and Osamyu Shinomura shared the 2008 Nobel prize in chemistry for

the discovery and exploitation of Cnidarian fluorescent proteins. Several further cycles of mutation, directed modification and evolutionary selection produced

mCherry, which is monomeric and has an

excitation maximum at 587nm and emission maximum at 610nm (3). The protein is widely used as a fluorescent tracer in transfection, transgenic, photobleaching and FRET type experiments. The prototype for these fluorescent proteins is Green Fluorescent Protein (GFP), which is a ~27kDa protein isolated originally from the jellyfish Aequoria victoria (4). The mCherry protein is similar in size and general structural properties to GFP (5,6), but, obviously, produces a red rather than a green fluorochrome. As with GFP, mCherry becomes fluorescent due to intrinsic properties requiring only molecular oxygen and so can be readily expressed in a

variety of systems.

Form: Liquid

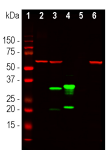
Background:

Buffer: Purified at 1mg/mL in 50% PBS, 50% glycerol, 5mM NaN3

Storage: Stable at 4°C for one year, for longer term store at -20°C

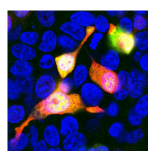


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Western blot analysis of HEK293 cell lysates, and recombinant protein solutions using mouse mAb to mCherry, dilution

1:1,000, in green [1] protein standard,
[2] HEK293, [3] HEK293 cells transfected
with mCherry-HA construct, [4] mCherry
recombinant protein, [5] GFP
recombinant protein, and [6] HEK293
transfected with GFP construct. Major
band at about 30kDa corresponds to
mCherry protein. mCherry antibody does
not react with GFP protein. The same
blot was simultaneously probed with
chicken pAb to HSP60, dilution 1:5,000
in red which reveals band at 60kDa seen
only in cell lysates.



Immunofluorescent analysis of HEK293 cells transfected with mCherry-HA, construct, in red, and stained with mouse mAb to mCherry, dilution 1:500, in green. The blue is Hoechst staining of nuclear DNA. mCherry antibody reveals mCherry protein expressed only in transfected cells which appear golden in color. Untransfected cells do not react with the antibody, as a result only their nuclei are visible.

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