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DATASHEET

Fluorescent Antibody Kit Atto565

Anti-Mouse IgG (H+L) Atto565

Goat-anti mouse IgG (H+L) Atto565

For Laboratory Use Only.

Not for Use in Diagnostic Processes.

Kit Content (Cat. #: 2107-1MG)

1.0mg Anti-Mouse IgG (H+L) Atto565 1.0ml Glycerol-PBS 50µg Mono-anti actin Product documentation & Certificate of Analysis

Product Documentation

Anti-Mouse IgG (H+L) Atto565

Anti-Mouse IgG (H+L) is an antigen-specific antibody. Affinity purification removed essentially all goat serum proteins, including immunoglobulins not specifically binding to mouse IgG. Goat anti-mouse IgG is conjugated to Atto565 (Abs.max. 563 nm; Em.max. 592 nm) and further purified by gel filtration.

Anti-Mouse IgG (H+L) Atto565 is supplied in unit sizes of 1.0mg.

Reconstitution of Antibodies with Glycerol-PBS

Add 0.5ml Glycerol-PBS to the lyophilized antibody to reconstitute a 2mg/ml stock solution. Vortex for 10sec until completely dissolved. Add 50 μ l Glycerol-PBS to the lyophilized primary antibody to reconstitute a 1mg/ml stock solution. Final concentrations of the antibody buffers: 0.01M sodium phosphate, 0.1M NaCl, pH 7.4, 5mM NaN₃ in 50% glycerol.

Working Dilution

Each individual user should determine the optimum working dilution empirically for the systems. Dilutions of 1:500 - 1:1500 are suitable for many applications.

Determining the Degree of Labeling (DOL)

1. Protein Concentration

Determination of the protein concentration by UV absorption measurement at 280nm ($\epsilon_{max} = 203,000 \, \text{M}^{-1} \text{cm}^{-1}$).

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2. Degree of Labelling

The degree of labeling (DOL or dye/protein ratio) is usually determined by absorption spectroscopy making use of the Lambert-Beer law: Absorbance (A) = extinction coefficient (\mathcal{E}) × molar concentration × path length (d). Simply measure the UV-VIS spectrum of the conjugate in solution in a quartz cuvette. Dilute the solution, if necessary to measure within the linear range.

$$DOL = \frac{A_{563} \cdot 203,000}{A_{280} - (A_{563} \cdot 0.26) \cdot 115,000}$$

 A_{563} = maximal absorbance at 563nm measured in a cuvette with a pathlength of 1 cm.

 A_{280} = maximal absorbance at 280nm measured in a cuvette with a pathlength of 1 cm.

203,000 = molar extinction coefficient (E) at the longest-wavelength absorption maximum (M⁻¹cm⁻¹).

120,000 = molar extinction coefficient (E) at the longest-wavelength absorption maximum (M-1cm-1).

0.16 = correction factor for the fluorophore's absorbance at 280nm.

Storage and Stability

For continuous use, store at 2-8 $^{\circ}$ C for up to three months. For extended storage, the solution may be frozen in working aliquots at -20 $^{\circ}$ C. Frozen aliquots are stable for at least six months. Avoid repeated freeze/thawing. If slight turbidity occurs upon prolonged storage, clarify the solution by centrifugation before use. Protect fluorescent conjugates from light.

Mono Anti-Actin

Monoclonal anti-actin (98% purity) recognizes skeletal and non-muscle actin isoforms. Isotype classified as an IgM, it reacts even stronger with Anti-Mouse IgG. In immunofluorescence microscopy samples are fixed with methanol to detect cytoplasmic actin, while fixation with para-formaldehyde leads to nuclear actin detection (Gonsior et al., 1999).

As immunogen for mono anti-actin a profilin-actin complex from calf thymus was used, and epitope mapping localized the following sequence (Gonsior et al.):

NVPAMYVAVLDSGVTHNVPIYHAIMRLDLA.

Mono anti-actin was tested on PtK2, SR-NRK, NRK-49F, L6 cells, C2C12, NIH-3T3, mouse myoblast and myotube cells.

Working Dilution

Each individual user should determine the optimum working dilution empirically for the systems. Dilutions of 1:100 – 1:300 with respect to the above mentioned fixation methods are sufficient for many applications.

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Reference:

Gonsior SM, et al.: Conformational difference between nuclear and cytoplasmic actin as detected by a monoclonal antibody. J Cell Sci 112, 797-809 (1999)

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