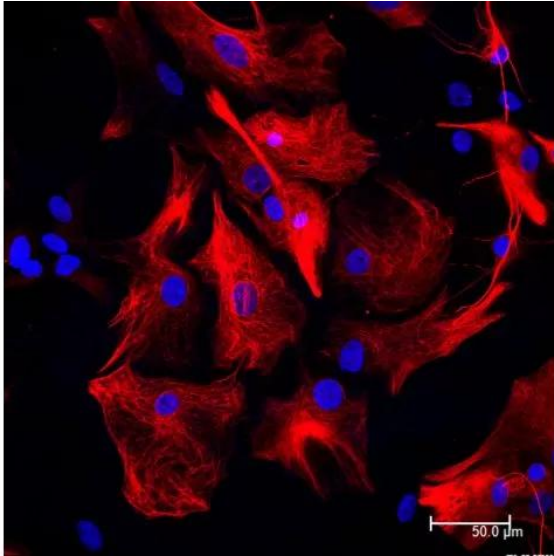




HORSE ANTI-RABBIT IGG ANTIBODY (H+L), DYLIGHT 594

SKU: DI-1094-1.5



DESCRIPTION

DyLight 594 Horse Anti-Rabbit IgG Antibody can be used for tissue staining and other applications. Optimal F/P ratios have been established for each conjugate to ensure maximum fluorescence with minimal background staining.

Features:

- Affinity-purified, ultrapure, high affinity antibody
- Thoroughly adsorbed against serum and immunoglobulins from potentially interfering species
- Recognizes both heavy and light chains (H+L)
- Optimally labeled with DyLight® 594 to provide the brightest label for fluorescence microscopy
- Supplied in solution
- Excitation: 592 nm
- Emission: 617 nm

For research use only. Not intended for therapeutic or diagnostic use in animals or humans.



- Color: Red

SPECIFICATIONS

Color of Fluorescence Red

Format Concentrate

Formulation 10 mM HEPES, 0.15 M NaCl, pH 7.5, 0.08% sodium azide.

Maximum Emission 617 nm

Maximum Excitation 592 nm

Unit Size 1.5 mg

Storage Instructions 2-8 °C

Usage Summary

Recommended concentration range for use 5-20 µg/ml. If this antibody is to be used in tissues which may contain cross-reacting endogenous immunoglobulins, dilution of this antibody may be made in buffers containing 2% normal serum from the same species as the tissue.

Applications

Immunofluorescence, In situ hybridization, Blotting Applications, Flow Cytometry/Cell Separation

Target Species Rabbit

Concentration 1.5 mg active conjugate/ml

Conjugate DyLight 594

Reactive Species Horse

Source Species Rabbit

Host Species Horse

TECHNICAL INFORMATION

The horse anti-rabbit Ig antibodies are prepared by hyperimmunizing animals in a manner that produces high affinity antibodies. These are then purified by an affinity chromatography procedure designed to remove any low affinity antibodies which may be present. Cross-reactivities that are likely to interfere with specific labeling are removed by solid-phase adsorption techniques. The final product is then subjected to rigorous quality control assays including immunodiffusion, solid-phase enzyme immunoassays, gel electrophoresis and solid-phase binding assays. In preparing the labeled antibodies, great care is taken to ensure the maximum degree of labeling with no alteration in the specificity and affinity of the antibody. The

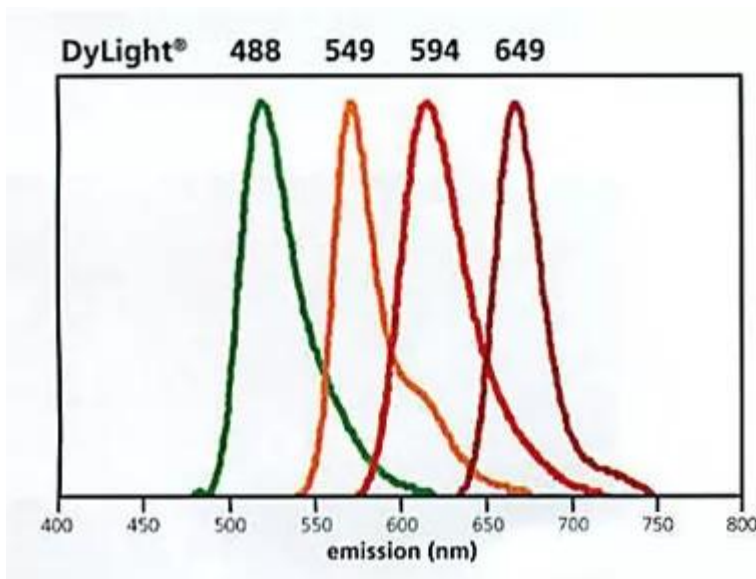
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labeled antibody then undergoes a further series of quality control assays, including immunohistochemical analysis.

DyLight™ fluorescent dyes are direct alternatives to traditional fluorophores such as fluorescein (FITC) and rhodamine. The excitation and emission spectra parallel that of other commercially available fluorescent reagents allowing for easy substitution into an existing protocol without requiring any further instrumentation or filter sets.

DyLight™ dyes offer a number of potential advantages including greater photostability and brighter fluorescence. DyLight™ dyes are completely stable over a pH range of 4-9 making them compatible with many aqueous-based buffers and diluents. The DyLight™ dyes can be applied as single labels or in combination with other DyLight™ dyes and fluorophores as part of a multiple immunofluorescent antigen staining methodology. The DyLight™ dyes currently offered are DyLight™ 488 (green), DyLight™ 549 (orange), DyLight™ 594 (red), and DyLight™ 649 (far red).



| Conjugate | Excitation maximum (nm) | Emission maximum (nm) | Spectrally similar dyes |
|-------------|-------------------------|-----------------------|-----------------------------|
| DyLight 488 | 493 | 518 | FITC, Alexa Fluor 488, Cy2 |
| DyLight 549 | 556 | 571 | TRITC, Alexa Fluor 555, Cy3 |
| DyLight 594 | 592 | 617 | Alexa Fluor 594, Texas Red |
| DyLight 649 | 655 | 670 | Alexa Fluor 647, Cy5 |

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CITATIONS

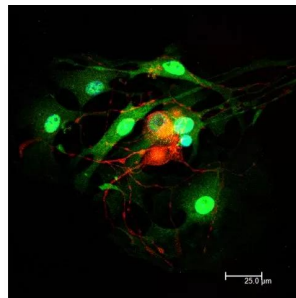
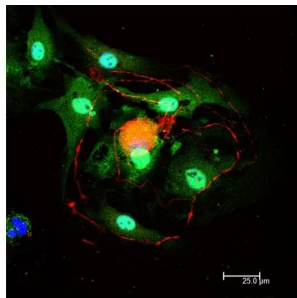
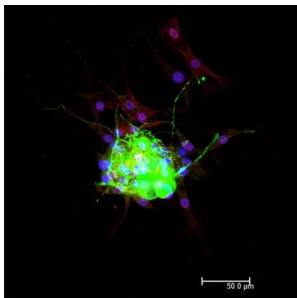
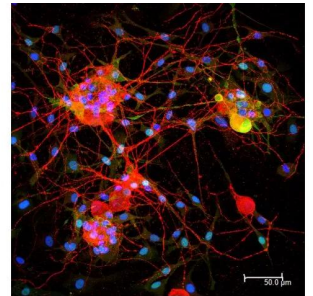
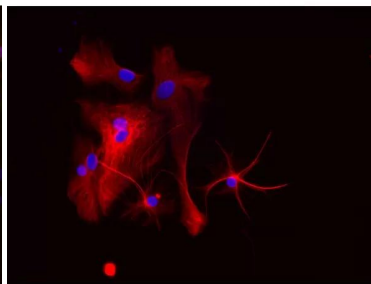
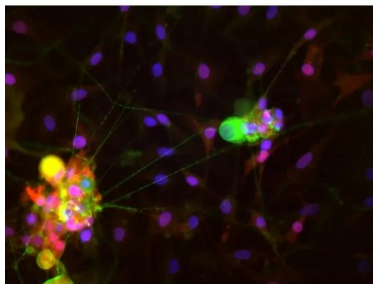
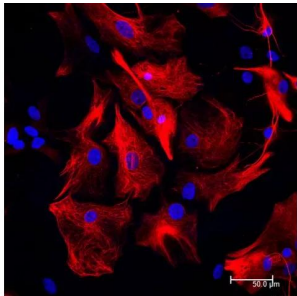


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DOCUMENTS

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