

# Rhodamine (TRITC)-conjugated Goat anti-Rabbit IgG (H+L)

Catalog No.: AS040 29 Publications

Basic Information	Background
Observed MW	Secondary antibodies are affinity-purified antibodies which will work with target- specific primary antibody in the detection, sorting or purification of its specified target.
Calculated MW	Secondary antibodies offer increased versatility enabling users to use many detection systems (e.g. HRP, AP, fluorescence). They can also provide greater sensitivity through
Category	signal amplification as multiple secondary antibodies . Most commonly, secondary
Secondary Antibody	antibodies are generated by immunizing the host animal (different from host species of primary antibody) with a pooled population of normal immunoglobulins from the host
Applications	species of primary antibody and can be further purified and modified (i.e. antibody
IHC-P,IF/ICC,FC	fragmentation, label conjugation, etc.) to ensure well-characterized specificity to corresponding normal immunoglobulins.
Cross-Reactivity	
Conjugate	
Rhodamine. Ex:550nm. Em:570nm.	

## **Recommended Dilutions**

# Immunogen Information

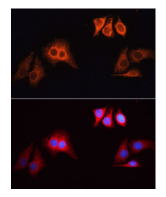
IHC-P	1:50 - 1:200	Gene ID	Swiss Prot
IF/ICC	1:50 - 1:200	Immunogen	
FC	1:50 - 1:200	Rabbit IgG <b>Synonyms</b>	
		Synonyms	

Contact		Product Information			
€	www.abclonal.com	<b>Source</b> Goat	<b>Isotype</b> TRITC conjugated IgG	<b>Purification</b> Affinity purification	
		Storage			

#### Storage

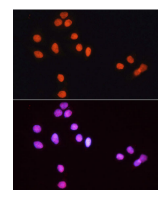
Store at -20°C. Avoid freeze / thaw cycles. Buffer: PBS with 0.025% Sodium Azide,0.75% BSA,50% glycerol,pH7.3.

## Validation Data



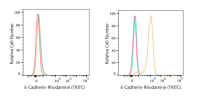
Immunofluorescence analysis of HeLa cells, using Rhodamine (TRITC) Goat Anti-Rabbit IgG (H+L) (A3716) as the primary antibody at dilution of 1:100. The cells were incubated with the primary antibody overnight at  $4^{\circ}$ C.

Secondary antibody: Rhodamine (TRITC) Goat Anti-Rabbit IgG (H+L) (AS040) at 1:200 dilution. Blue: DAPI for nuclear staining.



Immunofluorescence analysis of HeLa cells, using Rhodamine (TRITC) Goat Anti-Rabbit IgG (H+L) (A0942) as the primary antibody at dilution of 1:100. The cells were incubated with the primary antibody overnight at 4°C.

Secondary antibody: Rhodamine (TRITC) Goat Anti-Rabbit IgG (H+L) (AS040) at 1:100 dilution. Blue: DAPI for nuclear staining.



Flow cytometry: 1X10^6 K-562 cells (negative control,left) and A-431 cells (right) were surface-stained with Purified Rabbit anti-Human E-Cadherin mAb(5 µl/Test,orange line) or secondary antibody only (blue line). Non-fluorescently stained K-562 and A-431 cells were used as blank control (red line). Rhodamine (TRITC) Goat Anti-Rabbit IgG (H+L)(AS040, 1:200) was used as a secondary antibody.