

Datasheet for 100-4198**Biotin Antibody****Overview**

Description:	Anti-Biotin (RABBIT) Antibody - 100-4198
Item No.:	100-4198
Size:	2 mL
Applications:	ELISA, EM, IF, Multiplex, Other, WB
Reactivity:	Biotin
Host Species:	Rabbit

Product Details

Background:	Biotin Antibody detects Biotin. Biotin is a water-soluble B-complex vitamin (vitamin B7). It is composed of a ureido (tetrahydroimidizalone) ring fused with a tetrahydrothiophene ring. A valeric acid substituent is attached to one of the carbon atoms of the tetrahydrothiophene ring. Biotin is a coenzyme for carboxylase enzymes, involved in the synthesis of fatty acids, isoleucine, and valine, and in gluconeogenesis. Biotin is necessary for cell growth, the production of fatty acids, and the metabolism of fats and amino acids. Anti-Biotin Antibody is ideal for investigators involved in Cell Signaling and Cell Biology research.
Synonyms:	rabbit anti-biotin antibody, rabbit anti biotin
Host Species:	Rabbit
Clonality:	Polyclonal
Format:	Antiserum

Target Details

Reactivity:	Biotin
Immunogen Type:	Native Protein
Immunogen:	Biotin conjugated to Keyhole Limpet Hemocyanin (KLH)
Purity/Specificity:	This product was prepared from monospecific antiserum by a delipidation and defibrination. Assay by immunoelectrophoresis resulted in a single precipitin arc against anti-rabbit serum, biotinylated IgG and Biotinylated BSA.

Application Details

Tested Applications:	ELISA
Suggested Applications:	EM, IF, Multiplex, Other, WB (Based on references)
Application Note:	Anti-Biotin has been tested in ELISA and is suitable for immunoblotting (western or dot blot), immunoprecipitation and most immunological methods requiring high titer and specificity.
Assay Dilutions:	All assays should be optimized by the user. Recommended dilutions (if any) may be listed below.
ELISA:	1:100,000
IHC:	User Optimized
WB:	1:10,000 - 1:25,000

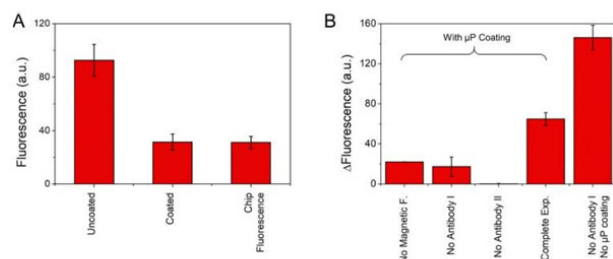
Formulation

Physical State:	Lyophilized
Concentration:	90 mg/mL by Refractometry
Buffer:	0.01 M Sodium Phosphate, 0.15 M Sodium Chloride, pH 7.2
Preservative:	0.01% (w/v) Sodium Azide
Stabilizer:	None
Reconstitution Volume:	2.0 mL
Reconstitution Buffer:	Restore with deionized water (or equivalent)

Shipping & Handling

Shipping Condition:	Ambient
Storage Condition:	Store vial at 4° C prior to restoration. For extended storage aliquot contents and freeze at -20° C or below. Avoid cycles of freezing and thawing. Centrifuge product if not completely clear after standing at room temperature. This product is stable for several weeks at 4° C as an undiluted liquid. Dilute only prior to immediate use.
Expiration:	Expiration date is one (1) year from date of receipt.

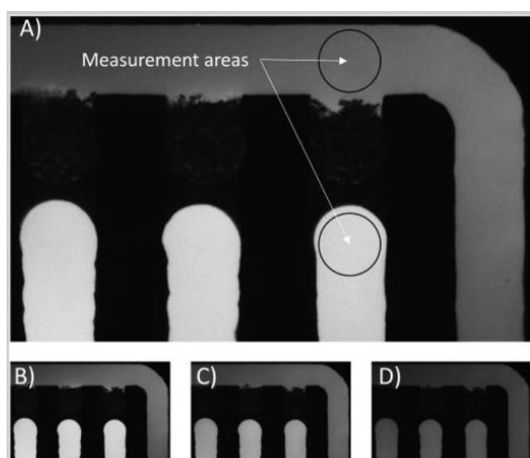
Images



Figure

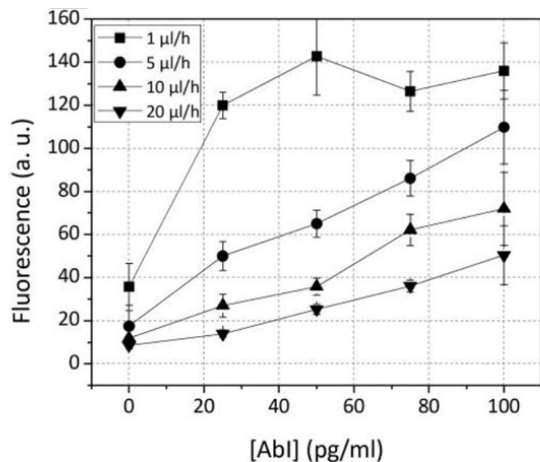
(a) Channels coating: remnant fluorescence due to nonspecific binding of anti-rabbit IgG labeled with B-rhodamine (p/n 611-1003) [AbII] at a concentration of 100 μ g/ml in uncoated and coated microchannels with silane-PEG.

(b) Fluorescence obtained from the complete immunoassay at a flow rate of 5 μ l/h and at an anti-biotin rabbit IgG (p/n 100-4198) [AbI] concentration of 50 pg/ml is compared to the fluorescence obtained from the immunoassay performed without applying the magnetic field (column 1), without adding anti-biotin rabbit IgG (p/n 100-4198) [AbI] (column 2), without adding anti-rabbit IgG labeled with B-rhodamine (p/n 611-1003) [AbII] (column 3), and, finally, the efficacy of the microparticles coating was tested by performing the immunoassay without anti-biotin rabbit IgG (p/n 100-4198) [AbI] and with noncoated microparticles (column 5). The level of fluorescence of the two first columns in b results from nonspecific interactions of anti-rabbit IgG labeled with B-rhodamine (p/n 611-1003) [AbII] with the microparticles. Error bars are standard deviation. FIG. 5. PMID: 32038740.



Figure

Typical experiment image where the complexes Ag–AbI–AbII are attached to the nanoparticles trapped in the magnetic traps and the fluorogenic substrate is flowed at different rates, where the measurement areas selected for the fluorescence analysis are shown. Measurement with a flow rate of (a) 1 μ l/h, (b) 2 μ l/h, (c) 5 μ l/h, and (d) 10 μ l/h. Experiment with anti-rabbit IgG labeled alkaline phosphatase was used (p/n 611-1502) and an anti-biotin rabbit IgG (p/n 100-4198) [AbI] concentration of 100 pg/ml. FIG. 6. PMID: 32038740.



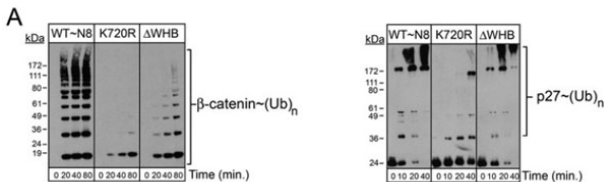
Figure

Shows the fluorescence difference from immunoassays at various concentrations of the primary antibody. Calibration curve: fluorescence difference from immunoassays with varying concentrations of anti-biotin rabbit IgG (p/n 100-4198) [Abl] at different flow rates of the fluorogenic enzyme substrate anti-rabbit IgG labeled alkaline phosphatase was used (p/n 611-1502). Each experiment was repeated on three different devices. FIG. 7. PMID: 32038740.

Western Blot

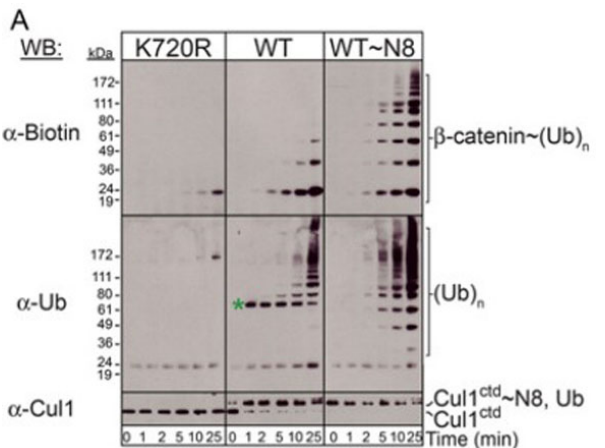
Western Blot of Anti-Biotin Antibody. Functional analysis of mutations influencing RING conformational flexibility.

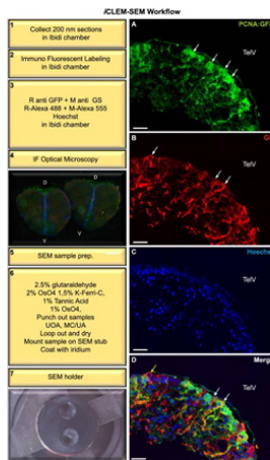
(A) Time-courses of polyubiquitination by SCFs reconstituted with fully-NEDD8ylated wild-type Cul1-Rbx1 (WT~N8), the non-NEDD8ylatable control (K720R), and the Cul1-Rbx1 WHB deletion mutant (Δ WHB). Left - SCF β TRCP-mediated polyubiquitination of a biotin-labeled β -catenin phosphopeptide, detected by western blotting with anti-biotin antisera. Right - SCFSkp2/CksHs1-mediated polyubiquitination of phospho-p27, detected by western blotting with anti-p27 antisera. Figure 5. PMID: 18805092.



Western Blot

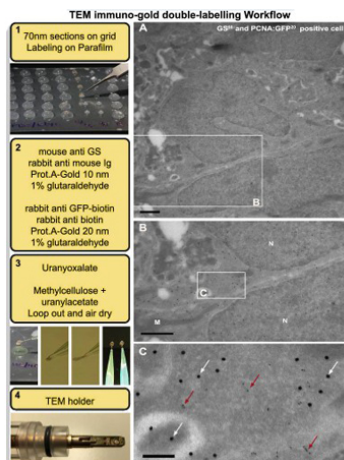
Western Blot of Anti-Biotin antibody. Conformational control of CRL activities. (A) Polyubiquitination reactions with SCF β TRCP/ β -catenin phosphopeptide (left), and SCFSkp2/CksHs1/ phospho-p27 (right) reconstituted with non-NEDD8ylatable (K720R), un-NEDD8ylated wild-type (WT), and fully NEDD8ylated Cul1-Rbx1 (WT~N8). Reaction products were detected by immunoblotting, top panels with anti-biotin (left) or anti-p27 (right), middle with anti-His (Ubiquitin; green * - Cul1~Ubiquitin), and lower with anti-Cul1 C-terminus antisera. Figure 7. PMID: 18805092.





Immunofluorescence Microscopy

iCLEM-SEM workflow and immunofluorescent labelling on Tokuyasu chemical-fixed cryosections. Left: Major steps required for iCLEM-SEM workflow (steps 1–6). (A) Green Fluorescent Protein (GFP)-positive labelling (green) of proliferating cells from the Tg(PCNA:GFP) transgenic line (white arrows). (B) Glutamine Synthetase (GS)-positive labelling (red) of glial cells (white arrows). (C) Hoechst counterstaining (blue). (D) Multi-channel merge showing co-labelling of PCNA:GFP-positive cells with the glial marker (orange cells; white arrows) in the cytoplasm of a small number of cells. Differences in the intensity of antibody staining can additionally be noted (compare white arrows and yellow arrow). D dorsal; V ventral. TelV telencephalic ventricle. Scale bar (A–D) 20 µm. Fig 2. PMID: 33441723



Immunofluorescence Microscopy

TEM immuno-gold double-labelling workflow and example of positive labelling on Tokuyasu chemical-fixed cryosections. Left: Major steps (1–4) required for TEM immuno-gold double-labelling workflow and associated images. (A,B,C) Cytosolic PCNA:GFP+/GS+ immuno-gold double-labelling at low (A) and high (B,C) magnifications demonstrating co-localization of the 10 nm (GS; red arrowheads) and 20 nm (PCNA:GFP; white arrowheads) particles. N, nucleus. Scale bar (A) 2 µm; (B,C) 200 nm. Fig 6. PMID: 33441723

References

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