Prion Protein(PrP) Rabbit mAb

Catalog No: #48939

Package Size: #48939-1 50ul #48939-2 100ul



Orders: order@signalwayantibody.com Support: tech@signalwayantibody.com

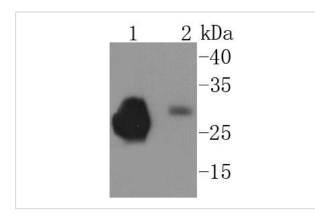
Description	
Product Name	Prion Protein(PrP) Rabbit mAb
Host Species	Recombinant Rabbit
Clonality	Monoclonal antibody
Clone No.	SC57-05
Purification	ProA affinity purified
Applications	WB, ICC/IF, IHC, FC
Species Reactivity	Hu, Ms, Rt
Immunogen Description	recombinant protein
Other Names	Alternative prion protein; major prion protein antibody AltPrP antibody ASCR antibody CD230 antibody CD230
	antigen antibody CJD antibody GSS antibody KURU antibody Major prion protein antibody p27 30 antibody
	PRIO_HUMAN antibody Prion protein antibody Prion related protein antibody PRIP antibody PRNP antibody
	PrP antibody PrP27 30 antibody PrP27-30 antibody PrP33-35C antibody PrPC antibody PrPSc antibody Sinc
	antibody
Accession No.	Swiss-Prot#:P04156
Uniprot	P04156
GenelD	5621;
Calculated MW	28 kDa
Formulation	1*TBS (pH7.4), 1%BSA, 40%Glycerol. Preservative: 0.05% Sodium Azide.
Storage	Store at -20°C

Application Details

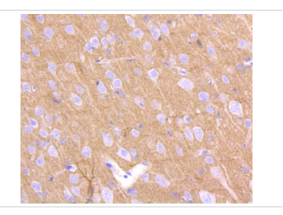
WB: 1:1,000-5,000IHC: 1:50-1:200

ICC: 1:50-1:200FC: 1:50-1:100

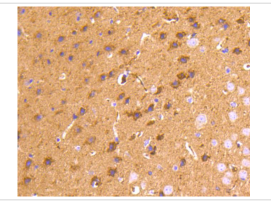
Images



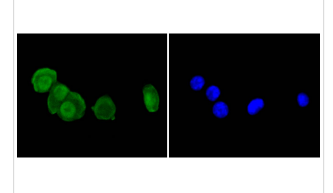
Western blot analysis of PrP on different lysates using anti-PrP antibody at 1/1,000 dilution. Positive control: Lane 1: Rat brain Lane 2: Mouse brain



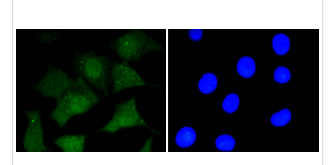
Immunohistochemical analysis of paraffin-embedded rat brain tissue using anti-PrP antibody. Counter stained with hematoxylin.



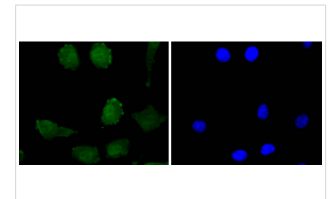
Immunohistochemical analysis of paraffin-embedded mouse brain tissue using anti-PrP antibody. Counter stained with hematoxylin.



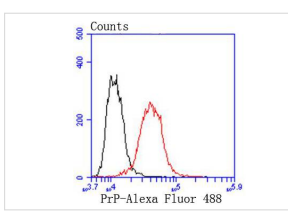
ICC staining PrP in N2A cells (green). The nuclear counter stain is DAPI (blue). Cells were fixed in paraformaldehyde, permeabilised with 0.25% Triton X100/PBS.



ICC staining PrP in SHG-44 cells (green). The nuclear counter stain is DAPI (blue). Cells were fixed in paraformaldehyde, permeabilised with 0.25% Triton X100/PBS.



ICC staining PrP in SH-SY-5Y cells (green). The nuclear counter stain is DAPI (blue). Cells were fixed in paraformaldehyde, permeabilised with 0.25% Triton X100/PBS.



Flow cytometric analysis of SH-SY-5Y cells with PrP antibody at 1/50 dilution (red) compared with an unlabelled control (cells without incubation with primary antibody; black). Alexa Fluor 488-conjugated goat anti rabbit IgG was used as the secondary antibody.

Background

Prion diseases, or transmissible spongiform encephalopathies (TSEs), are manifested as genetic, infectious or sporadic, lethal neurodegenerative disorders involving alterations of the prion protein (PrP). Characteristic of prion diseases, cellular PrP (PrPc) is converted to the disease form, PrPSc, through alterations in the protein folding conformations. PrPc is constitutively expressed in normal adult brain and is sensitive to proteinase K digestion, while the altered PrPSc conformation is resistant to proteases, resulting in a distinct molecular mass after PK treatment. Consistent with the transient infection process of prion diseases, incubation of PrPc with PrPSc both in vitro and in vivo produces PrPc that is resistant to protease degradation. Infectious PrPSc is found at high levels in the brains of animals affected by TSEs, including scrapie in sheep, BSE in cattle and Cruetzfeldt-Jakob disease in humans.

References

Note: This product is for in vitro research use only