## For Research Use Only

## Na 5-Hydroxymethylcytidine Monoclonal antibody, PBS Only

Catalog Number:68579-1-PBS

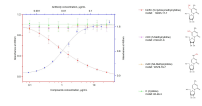


Basic Information	Catalog Number: 68579-1-PBS Size: 1 mg/ml Source: Mouse Isotype: IgG1	GenBank Accession Number: GeneID (NCBI): Full Name:	Purification Method: Protein G purification CloneNo.: 1E6C6
Applications	Tested Applications: Indirect ELISA,ELISA,Dot Blot Species Specificity: Human		
Background Information	Oxidation of 5-methylcytosine in DNA by ten-eleven translocation (Tet) family of enzymes has been demonstrated to play a significant role in epigenetic regulation in mammals. Recent reaearch shows that Tet enzymes also possess the activity of catalyzing the formation of 5-hydroxymethylcytidine (5-hmrC) in RNA. It is known that RNA carries more than 100 distinct types of modifications, and these modifications modulate the structure and functions of RNA. Ribonucleoside 5-methylcytidine (m5C) is subject to oxidative processing in mammals, forming 5-hydroxymethylcytidine (hm5C) and 5-formylcytidine (f5C). Researchers have identified hm5C in total RNA from all three domains of life and in polyA-enriched RNA fractions from mammalian cells. This suggests m5C oxidation is a conserved process that could have critical regulatory functions inside cells (PMID: 25676849).		
Storage	Storage: Store at -80°C. The product is shipped with ice packs Storage Buffer: PBS Only	s. Upon receipt, store it immediately at	t-80°C

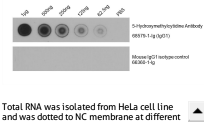
For technical support and original validation data for this product please contact:T: 4006900926E: Proteintech-CN@ptglab.comW: ptgcn.com

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## Selected Validation Data



Indirect ELISA and competitive ELISA results show that this antibody is specific to 5-Hydroxymethylcytidine (hm5C). Indirect ELISA was performed by coating BSA conjugated 5-Hydroxymethylcytidine (hm5C) at 10ng/well followed by blocking with 1% BSA. Serial diluted primary antibody was added to the plates and incubated at 37°C. HRP-goat anti-mouse was used for detection. Competitive ELISA was performed similarly except that different concentration of 5-



and was dotted to NC membrane at different amount as indicated above the dots. The membrane was blocked with BSA and blotted with 5-Hydroxymethylcytidine (hm5C) antibody 68579-1-1g at 1:5000 followed by incubation of HRP-goat anti-mouse secondary antibody. Signal was developed by ECL substrate. A parallel dot blot was performed using mouse IgG1 isotype control antibody 66360-1-1g at the same dose. This data was