For Research Use Only

Pseudouridine Monoclonal antibody

Catalog Number:68578-1-lg



Basic Information	Catalog Number:	GenBank Accession Number:	Purification Method:
	68578-1-lg Size: 150ul , Concentration: 1000 ug/ml by Nanodrop; Source: Mouse Isotype: IgG1	GeneID (NCBI): Full Name:	Protein G purification CloneNo.: 1G7C7 Recommended Dilutions: ELISA 1:2000-1:20000 Dot Blot 1:1000-1:4000
Applications	Tested Applications: Dot Blot, ELISA Species Specificity: chemical compound	Positive Co ELISA : Pset Dot Blot : RI	udouridine,
Background Information	Pseudouridine is an isomer of the nucleoside uridine in which the uracil is attached via a carbon-carbon instead of a nitrogen-carbon glycosidic bond. Pseudouridine is the most abundant RNA modification in cellular RNA (PMID: 29104216). After transcription and following synthesis, RNA can be modified with over 100 chemically distinct modifications. These can potentially regulate RNA expression post-transcriptionally, in addition to the four standard nucleotides and play a variety of roles in the cell including translation, localization and stabilization of RNA. Pseudouridine in rRNA and tRNA has been shown to fine-tune and stabilize the regional structure and help maintain their functions in mRNA decoding, ribosome assembly, processing and translation (PMID: 10902565, PMID: 23391857, PMID: 28045575). Pseudouridine in snRNA has been shown to enhance spliceosomal RNA-pre-mRNA interaction to facilitate splicing regulation (PMID 27268497).		
Storage	Storage: Store at -20°C. Stable for one year aft Storage Buffer: PBS with 0.02% sodium azide and 50	% glycerol pH 7.3.	
*** 20ul sizes contain 0.1% BSA	Aliquoting is unnecessary for -20 $^\circ$ C s	torage	

For technical support and original validation data for this product please contact:T: 1 (888) 4PTGLAB (1-888-478-4522) (toll free
in USA), or 1(312) 455-8498 (outside USA)E: proteintech@ptglab.comW: ptglab.comW: ptglab.com

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Selected Validation Data



Indirect ELISA and competitive ELISA results show that this antibody is specific to Pseudouridine. Indirect ELISA was performed by coating BSA conjugated Pseudouridine 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 Pseudouridine or its structure analogue



Total RNA was isolated from HeLa cell line and was dotted to NC membrane at different amount as indicated above the dots. The membrane was blocked with BSA and blotted with Pseudouridine antibody 68578-1-1g at 1:2000 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.