SARS-CoV-2 Nucleocapsid Phosphoprotein Recombinant antibody

Catalog Number:80027-1-RR

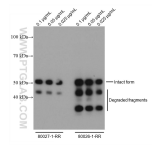
Basic Information	Catalog Number: 80027-1-RR	GenBank Accession Number: NC_045512	Purification Method: Protein A purification	
	100ul , Concentration: 1000 µg/ml by Nanodrop;	GeneID (NCBI): 43740575 Full Name: COVID-19 N Protein	CloneNo.: 8C20 Recommended Dilutions: WB 1:5000-1:50000	
	Immunogen Catalog Number: AG30676			
Applications	Tested Applications: WB,ELISA Species Specificity: virus		Positive Controls: WB : Eukaryotic nucleocapsid phosphoprotein, Ag30676	
Background Information	The nucleocapsid (N) protein has multiple functions including formation of nucleocapsids, signal transduction virus budding, RNA replication, and mRNA transcription. N protein is an important antigen for coronavirus, and it is normally highly conserved, with a molecular weight of about 50 kDa. it can be used as a marker in diagnostic assays due to its high immunogenicity (PMID: 32416961, PMID: 32235387). A sandwich ELISA for COVID-19 N Protein can be assembled by using 80027-1-RR as capture antibody and conjugated 80026-1-RR for detection.			
Storage	Storage: Store at -20°C. 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 E: proteintech@ptglab.com in USA), or 1(312) 455-8498 (outside USA) W: ptglab.com

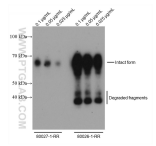
This product is exclusively available under Proteintech Group brand and is not available to purchase from any other manufacturer.



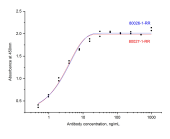
Selected Validation Data



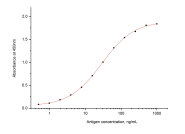
E.coli expressed SARS-CoV-2 Nucleocapsid Phosphoprotein (Cat.NO. Ag30676) was subjected to SDS-PAGE followed by western blot with 80027-1-RR and 80026-1-RR at various work concentration.



Eukaryotic expressed SARS-CoV-2 Nucleocapsid Phosphoprotein was subjected to SDS-PAGE followed by western blot with 80027-1-RR and 80026-1-RR at various work concentration.



Indirect ELISA was carried out by coating eukaryotic expressed N protein at 70 ng/well followed by blocking and adding serial diluted primary antibody 80026-1-RR and 80027-1-RR respectively. Signal was developed with TMB and stopped by H2SO4. Signal strength was measured by absorbance at 450 nm.



Sandwich ELISA was carried out by coating 80027-1-RR at 80 ng/well followed by blocking and adding different concentration of eukaryotic expressed N protein (0.5-1000 ng/mL). HRPconjugated80026-1-RR was used at 1 µg/mL for detection. Signal was developed with TMB and stopped by H2SO4. Signal strength was measured by absorbance at 450 nm.