For Research Use Only

DFNA5/GSDME Polyclonal antibody

Catalog Number:30696-1-AP

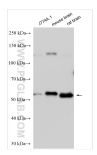


Basic Information	Catalog Number: 30696-1-AP	GenBank Accession Number: BC132303	Purification Method: Antigen affinity purification	
	Size: 300 µg/ml	GenelD (NCBI): 54722	Recommended Dilutions: WB 1:500-1:2000	
	Source: Rabbit	UNIPROT ID: Q9Z2D3		
	Isotype: IgG Immunogen Catalog Number: AG33622	Full Name: deafness, autosomal dominant 5 (human)		
		Calculated MW: 57 kDa		
		Observed MW: 57 kDa		
Applications	Tested Applications: Positive Controls:		Controls:	
	WB, ELISA Species Specificity: mouse	WB : J774A.1 cells, mouse brain tissue, rat brain tissue		
Background Information	DFNA5 (deafness, autosomal dominant 5), also known as GSDME or ICERE-1, is a 496 amino acid protein that is expressed in cochlea tissue, as well as in placenta, brain, heart, liver, lung and pancreas. Defects in the gene encoding DFNA5 are the cause of non-syndromic sensorineural deafness autosomal dominant type 5 (DFNA5), a form of sensorineural hearing loss that results from damage to one of various structures that receive sound information in the brain.			
Storage	Storage: Store at -20°C. Stable for one year after shipment. Storage Buffer: PBS with 0.02% sodium azide and 50% glycerol pH 7.3. Aliquoting is unnecessary for -20°C storage			

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



Various lysates were subjected to SDS PAGE followed by western blot with 30696-1-AP (DFNA5/GSDME antibody) at dilution of 1:1000 incubated at room temperature for 1.5 hours.