Datasheet for ABIN883796 anti-SMPD1 antibody (AA 201-300) (Cy5.5)

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Overview

Quantity:	100 μL
Target:	SMPD1
Binding Specificity:	AA 201-300
Reactivity:	Human, Mouse, Rat
Host:	Rabbit
Clonality:	Polyclonal
Conjugate:	This SMPD1 antibody is conjugated to Cy5.5
Application:	Western Blotting (WB), Immunofluorescence (Cultured Cells) (IF (cc)), Immunofluorescence (Paraffin-embedded Sections) (IF (p))

Product Details

Immunogen:	KLH conjugated synthetic peptide derived from human Acid sphingomyelinase
Isotype:	lgG
Cross-Reactivity:	Human, Mouse, Rat
Predicted Reactivity:	Dog,Cow,Pig,Rabbit
Purification:	Purified by Protein A.
Target Details	
Target:	SMPD1
Alternative Name:	Acid sphingomyelinase (SMPD1 Products)

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Background:	Synonyms: Acid sphingomyelinase, ASM, ASM_HUMAN, aSMase, NPD, Smpd1, Sphingomyelin	
	phosphodiesterase 1 acid lysosomal, Sphingomyelin phosphodiesterase.	
	Background: Converts sphingomyelin to ceramide. Also has phospholipase C activities toward	
	1,2-diacylglycerolphosphocholine and 1,2-diacylglycerolphosphoglycerol. Isoform 2 and	
	isoform 3 have lost catalytic activity.Involvement in disease: Defects in SMPD1 are the cause o	
	Niemann-Pick disease type A (NPDA) , also known as Niemann-Pick disease classical infantile	
	form. It is an early-onset lysosomal storage disorder caused by failure to hydrolyze	
	sphingomyelin to ceramide. It results in the accumulation of sphingomyelin and other	
	metabolically related lipids in reticuloendothelial and other cell types throughout the body,	
	leading to cell death. Niemann-Pick disease type A is a primarily neurodegenerative disorder	
	characterized by onset within the first year of life, mental retardation, digestive disorders, failure	
	to thrive, major hepatosplenomegaly, and severe neurologic symptoms. The severe	
	neurological disorders and pulmonary infections lead to an early death, often around the age of	
	four. Clinical features are variable. A phenotypic continuum exists between type A (basic	
	neurovisceral) and type B (purely visceral) forms of Niemann-Pick disease, and the intermediate	
	types encompass a cluster of variants combining clinical features of both types A and B.	
Gene ID:	6609	
UniProt:	P17405	
Application Details		
Application Notes:	IF(IHC-P) 1:50-200	
	IF(IHC-F) 1:50-200	
	IF(ICC) 1:50-200	
Restrictions:	For Research Use only	
Handling		
Format:	Liquid	
Concentration:	1 μg/μL	
Buffer:	Aqueous buffered solution containing 0.01M TBS (pH 7.4) with 1 % BSA, 0.03 % Proclin300 ar 50 % Glycerol.	
Preservative:	ProClin	

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	handled by trained staff only.
Storage:	-20 °C
Storage Comment:	Store at -20°C. Aliquot into multiple vials to avoid repeated freeze-thaw cycles.
Expiry Date:	12 months