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Datasheet for ABIN914313 anti-SAMHD1 antibody (AA 256-370) (Biotin)



Overview

Quantity:	100 µL
Target:	SAMHD1
Binding Specificity:	AA 256-370
Reactivity:	Mouse
Host:	Rabbit
Clonality:	Polyclonal
Conjugate:	This SAMHD1 antibody is conjugated to Biotin
Application:	Western Blotting (WB), ELISA, Immunohistochemistry (Paraffin-embedded Sections) (IHC (p)), Immunohistochemistry (Frozen Sections) (IHC (fro))

Product Details

Immunogen:	KLH conjugated synthetic peptide derived from human SAMHD1
lsotype:	lgG
Cross-Reactivity:	Mouse
Predicted Reactivity:	Human,Rat,Cow,Sheep,Horse
Purification:	Purified by Protein A.
Target Details	
Target:	SAMHD1

Target:	SAMHD1
Alternative Name:	SAMHD1 (SAMHD1 Products)

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Target Details	
Target Details Background:	Synonyms: DCIP, Dendritic cell derived NG induced protein, Dendritic cell-derived NG-induced protein, HD domain containing 1, HDDC1, Mg11, Monocyte protein 5, MOP 5, MOP5, OTTHUMP00000030889, SAM domain and HD domain 1, SAM domain and HD domain containing protein 1, SAM domain and HD domain-containing protein 1, SAMH1_HUMAN, Samhd1, SBBI88. Background: Putative nuclease involved in innate immune response by acting as a negative regulator of the cell-intrinsic antiviral response. May play a role in mediating proinflammatory responses to TNF-alpha signaling.Tissue specificity:Expressed in heart, skeletal muscle, spleen, liver, small intestine, placenta, lung and peripheral blood leukocytes. No expression is seen in brain and thymus.Involvement in disease:Defects in SAMHD1 are the cause of Aicardi-Goutieres syndrome type 5 (AGS5). A form of Aicardi-Goutieres syndrome, a genetically heterogeneous disease characterized by cerebral atrophy, leukoencephalopathy, intracranial calcifications, chronic cerebrospinal fluid (CSF) lymphocytosis, increased CSF alpha-interferon, and negative serologic investigations for common prenatal infection. Clinical features as thrombocytopenia, hepatosplenomegaly and elevated hepatic transaminases along with intermittent fever may erroneously suggest an infective process. Severe neurological dysfunctions manifest in infancy as progressive microcephaly, spasticity, dystonic posturing
	and profound psychomotor retardation. Death often occurs in early childhood.
Gene ID:	4861
UniProt:	Q9Y3Z3

Application Details

Application Notes:	WB 1:300-5000
	IHC-P 1:200-400
	IHC-F 1:100-500
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 and 50 % Glycerol.
Preservative:	ProClin

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Handling	
Precaution of Use:	This product contains ProClin: a POISONOUS AND HAZARDOUS SUBSTANCE, which should be handled by trained staff only.
Storage:	-20 °C
Storage Comment:	Store at -20°C for 12 months.
Expiry Date:	12 months