

ARG54718 anti-SIRT1 antibody [1089CT5.3.1]

Package: 100 μl Store at: -20°C

Summary

Product Description	Mouse Monoclonal antibody recognizes SIRT1
Tested Reactivity	Hu, Ms
Tested Application	WB
Host	Mouse
Clonality	Monoclonal
Clone	1089CT5.3.1
Isotype	IgG3, kappa
Target Name	SIRT1
Species	Human
Immunogen	Synthetic peptide corresponding to aa. 1-180 of Human SIRT1 (NP_001135970.1).
Conjugation	Un-conjugated
Alternate Names	75SirT1; SIR2L1; SIR2alpha; SIR2-like protein 1; EC 3.5.1; NAD-dependent protein deacetylase sirtuin-1; SIR2; hSIRT1; Regulatory protein SIR2 homolog 1; hSIR2

Application Instructions

Application table	Application	Dilution
	WB	1:1000
Application Note	* The dilutions indicate recommended starting dilutions and the optimal dilutions or concentrations should be determined by the scientist.	
Positive Control	A549	

Properties

Purification	Protein G purified
Buffer	PBS and 0.09% (W/V) Sodium azide
Preservative	0.09% (W/V) Sodium azide
Storage instruction	For continuous use, store undiluted antibody at 2-8°C for up to a week. For long-term storage, aliquot and store at -20°C or below. Storage in frost free freezers is not recommended. Avoid repeated freeze/thaw cycles. Suggest spin the vial prior to opening. The antibody solution should be gently mixed before use.
Note	For laboratory research only, not for drug, diagnostic or other use.

Bioinformation

Database links	GenelD: 23411 Human
	GeneID: 93759 Mouse
	Swiss-port # Q923E4 Mouse
	Swiss-port # Q96EB6 Human
Gene Symbol	SIRT1
Gene Full Name	sirtuin 1
Background	This gene encodes a member of the sirtuin family of proteins, homologs to the yeast Sir2 protein. Members of the sirtuin family are characterized by a sirtuin core domain and grouped into four classes. The functions of human sirtuins have not yet been determined; however, yeast sirtuin proteins are known to regulate epigenetic gene silencing and suppress recombination of rDNA. Studies suggest that the human sirtuins may function as intracellular regulatory proteins with mono-ADP-ribosyltransferase activity. The protein encoded by this gene is included in class I of the sirtuin family. Alternative splicing results in multiple transcript variants. [provided by RefSeq, Dec 2008]
Function	NAD-dependent protein deacetylase that links transcriptional regulation directly to intracellular energetics and participates in the coordination of several separated cellular functions such as cell cycle, response to DNA damage, metobolism, apoptosis and autophagu. Can modulate chromatin function through deacetylation of histones and can promote alterations in the methylation of histones and DNA, leading to transcriptional repression. Deacetylates a broad range of transcription factors and corregulators, thereby regulating target gene expression positively and negatively. Serves as a sensor of the cytosolic ratio of NAD(+)/NADH which is altered by glucose deprivation and metabolic changes associated with caloric restriction. Is essential in skeletal muscle cell differentiation which also involves 5'-MAP-activated protein kinase (AMPK) and nicotinamide phosphoribosyltransferase (NAMPT). Component of the eNSC Complex is able to sense the energy status and acts by recruiting histone-modifying enzymes. The eNoSC Complex is able to sense the energy status of cell: upon glucose starvation, elevation of NAD(+)/NADP(-) ratio activates SIRTI, leading to histone H3 deacetylation followed by dimethylation of H3 at 'lys-9' (H3K9me2) by SUV39H1 and the formation of slient chromatin in the rDNA locus. Deacetylates 'Iys-16' of bistone H4 (in vitro). Involved in NR082/SHP corepression function through chromatin remodeling: Recortlet to LRH1 target gene promoters by NR082/SHP thereby stimulating histone H3 and H4 deacetylation leading to transcriptional repression. Proposed to contribute to genomic integrity via positive regulation of tel meres durages 20V39H1 (D9V39H1 polyubiquitination by MDM2. This increase in SUV39H1 West SH3H1 target gene locustative thereby SH3H1 throver in CH, which in turn seems to accelerate renewal of the heterochromatin which correlates with greater genomic integrity during stress response. Deacetylates 'US-382' of p53/TP53 and impairs its ability to induce transcription-adtry transcription at thry:

	mechanism after reach round of transcription is proposed. Involved in lipid metabolism. Implicated in regulation of adjogenesis and fat mobilization in white adjocyces by repression of PPARG which probably involves association with NCOR1 and SMRT/NCOR2. Deacetylates ACSS2 leading to its activation of PPARGC1A is required to activate fatty acid oxidation in skeltel muscle under low-glucose conditions and is involved in gulucose homeostasis. Involved in regulation of PPARA and fatty acid beta- coidation in liver. Involved in positive regulation of insulin secretion in pancreatic beta cells in response to glucose; the function seems to imply transcriptional repression of UCP2. Proposed to deacetylate IRS2 thereby facilitating its insulin-induced tyrosine phosphorylation. Deacetylates SREPE1 isoform SREPP-1C thereby decreasing its stability and transactivation in lipogenic gene expression. Involved in DNA damage response by repressing genes which are involved in DNA repair and SIRT1-deacetylated XPA interacts with RPA2. Also involved in DNA repair DNA double-strain breaks by homologous recombination and specifically single-strand annealing independently of XRCC6/Ku70 and NBN. Transcriptional suppression of XPC probably involves 24.4RBL2 suppressor complex and protein kinase B (ART) signaling. Transcriptional suppression of TP23 probably involves 2EP4 and PCAF. Deacetylates WRN thereby regulating its helicase and exonuclease activities and regulates WRN nuclear transforation in response to DNA damage. Deacetylates XPA1 at 11, VeS-6 and 11, xr-7 and stimulates cellular AP endonuclease activity by promoting the association of AFX1 to XRCC1. Increases p33/TP53-mediated transcription-independent apoptosis by blocking nuclear translocatof, with Nuclear transformic p53/TP53 and probably redirecting it to mitochondria thereby inhibiting stress- induced apoptosis. Is involved in a nothenypacy, resumably by deacetylating ATGS, ATG7 and MAP1L3B/ATG8. Deacetylates XKT1 which leads to enhanced binding of AKT1
	PubMed:19220062, PubMed:19364925, PubMed:19690166, PubMed:19934257, PubMed:20097625,
Research Area	Cell Biology and Cellular Response antibody; Cell Death antibody; Controls and Markers antibody; Gene Regulation antibody; Metabolism antibody; Microbiology and Infectious Disease antibody
Calculated Mw	82 kDa
РТМ	Methylated on multiple lysine residues; methylation is enhanced after DNA damage and is dispensable for deacetylase activity toward p53/TP53. Phosphorylated. Phosphorylated by STK4/MST1, resulting in inhibition of SIRT1-mediated p53/TP53

deacetylation. Phosphorylation by MAPK8/JNK1 at Ser-27, Ser-47, and Thr-530 leads to increased
nuclear localization and enzymatic activity. Phosphorylation at Thr-530 by DYRK1A and DYRK3 activates
deacetylase activity and promotes cell survival. Phosphorylation by mammalian target of rapamycin
complex 1 (mTORC1) at Ser-47 inhibits deacetylation activity. Phosphorylated by CaMK2, leading to
increased p53/TP53 and NF-kappa-B p65/RELA deacetylation activity (By similarity). Phosphorylation at
Ser-27 implicating MAPK9 is linked to protein stability. There is some ambiguity for some phosphosites:
Ser-159/Ser-162 and Thr-544/Ser-545.
Proteolytically cleaved by cathepsin B upon TNF-alpha treatment to yield catalytic inactive but stable
SirtT1 75 kDa fragment (75SirT1).
S-nitrosylated by GAPDH, leading to inhibit the NAD-dependent protein deacetylase activity.Cellular LocalizationCytoplasm, Mitochondrion, Nucleus

Images

