Revision 1

Solution of the second	Histone H2 XP [®] Rabbi ⁻	B (Lys15) t mAb				CHNOLOGY
Store					Orders:	877-616-CELL (2355) orders@cellsignal.com
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	nhy Not for Lloo i	n Diagnostia Drog	oduroo	3 Trask L	ane Danvers Ma	ssachusetts 01923 USA
For Research Use O	Reactivity:	Sensitivity:	MW (kDa):	Source/Isotype:	UniProt ID:	Entrez-Gene Id:
WB, IP, IF-IC, ChIP	H M R Mk	Endogenous	14	Rabbit IgG	#P33778	3018
Product Usage Information				body and 10 µg of chro impleChIP [®] Enzymatic		ely 4 x 10 ⁶ cells) per IP.
	4	pplication				Dilution
	V	Vestern Blotting				1:1000
	Ir	nmunoprecipitation				1:50
	Ir	mmunofluorescence (Immunocytochem	istry)		1:1600
	C	Chromatin IP				1:50
Storage				.5), 150 mM NaCl, 100 not aliquot the antibody		ycerol and less than
Specificity / Sei	or			Rabbit mAb recognizes ody does not cross-rea		of histone H2B protein acetylated at Lys5,
Species predict react based on sequence home	100%	ebrafish, Bovine, Pig,	Horse			
Source / Purific		onoclonal antibody is residues surrounding		0	synthetic acetylated	peptide corresponding
Background	bli be ac ac ar ne nu va th br du (a tra ch Ly re Ly (1 (1 ph ph at	bock of chromatin. Orig een shown to be dyna actylation, phosphoryl actylation, phosphoryl actyltransferases acet ad 20) at gene promot autralizes the positive acleosome interaction arious DNA-binding pr at facilitate recruitmer omodomain, which bi uring transcriptional ac lso known as RNF20/ anscribed region of ac aromatin remodeling (ars79, two additional hi sponse to metabolic s as36, both at promoter 1). In response to mu 2). Upon induction of hosphorylation of histo hosphorylated at irrad hosphorylation at Ser20/ anscribed region of action and the sponse to metabolic s as36, both at promoter and the sponse to mu and the spons	ginally thought to fi mic proteins, under ation, methylation, ylate multiple lysin ters during transcr charge of these d s, thereby destabi- oteins (4,5). In addi- to f many transcri- nds to acetylated crivation by the RA (RNF40) (7). Mono- trive genes and sti 7-9). In addition, it stone modification stress, AMPK is re rs and in transcribu- litiple apoptotic stir apoptosis, Mst1 is one H2B during ch iation-induced DN L4 is rapid, depend	ergoing multiple types of and ubiquitination (1,2 are residues in the amino- iptional activation (1-3) omains and is believed lizing chromatin structu dition, acetylation of sp ption and chromatin re- lysine residues (6). His ND6 E2 protein in conju o-ubiquitinated histone	fold for DNA packag of post-translational b). The p300/CBP his o terminal tail of hist to terminal tail of hist to weaken histone- re and increasing th ecific lysine residue: gulatory proteins that tone H2B is mono-u nction with the BRE H2B Lys120 is asso elongation by facilit uent methylation of ptional initiation and enes and phosphory and may regulate tran osphorylated at Ser by caspase-3, lead Interestingly, histon e embryonic fibrobla tion of H2AX Ser13	ing, histones have now modifications, including stone one H2B (Lys5, 12, 15, of the histone tails DNA and nucleosome- ne access of DNA to s creates docking sites at contain a biquitinated at Lys120 1A/BRE1B E3 ligase ciated with the ating FACT-dependent histone H3 Lys4 and l elongation (10). In ylates histone H2B at ascriptional elongation 14 by the Mst1 kinase ing to global e H2B is rapidly asts (13). In this case, 9, and occurs in the
D		pair and apoptosis.		0		
Background Re	eterences 1.	Peterson, C.L. and L	aniel, M.A. (2004)	Curr Biol 14, R546-51.		

/24, 12:20 PM Acety	 I-Histone H2B (Lys15) (D8H1) XP® Rabbit mAb (#9083) Datasheet Without Images Cell Signaling ⁻ 2. Jaskelioff, M. and Peterson, C.L. (2003) Nat Cell Biol 5, 395-9. 3. Roth, S.Y. et al. (2001) Annu Rev Biochem 70, 81-120. 4. Workman, J.L. and Kingston, R.E. (1998) Annu Rev Biochem 67, 545-79. 5. Hansen, J.C. et al. (1998) Biochemistry 37, 17637-41. 6. Yang, X.J. (2004) Bioessays 26, 1076-87. 7. Kim, J. et al. (2009) Cell 137, 459-71. 8. Minsky, N. et al. (2008) Nat Cell Biol 10, 483-8. 9. Pavri, R. et al. (2006) Cell 125, 703-17. 10. Shilatifard, A. (2006) Annu Rev Biochem 75, 243-69. 11. Bungard, D. et al. (2003) Cell 113, 507-17. 13. Fernandez-Capetillo, O. et al. (2004) J Exp Med 199, 1671-7.
Species Reactivity	Species reactivity is determined by testing in at least one approved application (e.g., western blot).
Western Blot Buffer	IMPORTANT: For western blots, incubate membrane with diluted primary antibody in 5% w/v BSA, 1X TBS, 0.1% Tween® 20 at 4°C with gentle shaking, overnight.
Applications Key	WB: Western Blotting IP: Immunoprecipitation IF-IC: Immunofluorescence (Immunocytochemistry) ChIP: Chromatin IP
Cross-Reactivity Key	 H: human M: mouse R: rat Hm: hamster Mk: monkey Vir: virus Mi: mink C: chicken Dm: D. melanogaster X: Xenopus Z: zebrafish B: bovine Dg: dog Pg: pig Sc: S. cerevisiae Ce: C. elegans Hr: horse GP: Guinea Pig Rab: rabbit All: all species expected
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