#4030

Phospho-p53 (Ser15) (16G8) Mouse mAb (Biotinylated)



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For Research Use Only. Not for Use in Diagnostic Procedures.

Applications: W	Reactivity: H	Sensitivity: Endogenous	MW (kDa): 53	Source/Isotype: Mouse IgG1	UniProt ID: #P04637	Entrez-Gene Id: 7157
Product Usage Information		Application Western Blotting			Dilution 1:1000	
Storage				nM sodium phosphate (j d 50% glycerol. Store at -		
Specificity/Sens	sitivity			(Biotinylated) detects en pes not cross-react with		
Source / Purific	ation			unizing animals with a s er15 of human p53 prot		eptide
Description		This Cell Signaling Tec	hnology (CST) antib	ody is conjugated to bic	tin under optimal c	conditions.
Background		The p53 tumor suppressor protein plays a major role in cellular response to DNA damage and other genomic aberrations. Activation of p53 can lead to either cell cycle arrest and DNA repair or apoptosis (1). p53 is phosphorylated at multiple sites <i>in vivo</i> and by several different protein kinases <i>in vitro</i> (2,3). DNA damage induces phosphorylation of p53 at Ser15 and Ser20 and leads to a reduced interaction between p53 and its negative regulator, the oncoprotein MDM2 (4). MDM2 inhibits p53 accumulation by targeting it for ubiquitination and proteasomal degradation (5,6). p53 can be phosphorylated by ATM, ATR, and DNA-PK at Ser15 and Ser37. Phosphorylation impairs the ability of MDM2 to bind p53, promoting both the accumulation and activation of p53 in response to DNA damage (4,7). Chk2 and Chk1 can phosphorylate p53 at Ser20, enhancing its tetramerization, stability, and activity (8,9). p53 is phosphorylated at Ser392 <i>in vivo</i> (10,11) and by CAK <i>in vitro</i> (11). Phosphorylation of p53 at Ser392 is increased in human tumors (12) and has been reported to influence the growth suppressor function, DNA binding, and transcriptional activation of p53 is mediated by p300 and CBP acetyltransferases. Inhibition of deacetylation suppressing MDM2 from recruiting HDAC1 complex by p19 (ARF) stabilizes p53. Acetylation appears to play a positive role in the accumulation of p53 protein in stress response (17). Following DNA damage, human p53 becomes acetylated at Lys382 (Lys379 in mouse) <i>in vivo</i> to enhance p53-DNA binding (18). Deacetylation of p53 occurs through interaction with the SIRT1 protein, a deacetylase that may be involved in cellular aging and the DNA damage response (19).				
Background Re	ferences	6. Honda, R. et al. (199 7. Tibbetts, R.S. et al. (8. Shieh, S.Y. et al. (199 9. Hirao, A. et al. (2000 10. Hao, M. et al. (1997) 11. Lu, H. et al. (1997) 12. Ullrich, S.J. et al. (1 13. Kohn, K.W. (1999)	emin Cancer Biol 5, (1997) Life Sci 60, 1 (1997) Life Sci 60, 1 (1999) Proc Natl Acc (1999) Genes Dev 1 (1999) Genes Dev 1 (1993) Proc Natl Acc (1997) Oncogene (1997) Oncogene (1997) Jon 131-40 (1998) Genes Dev	-11. ad Sci U S A 96, 13777-82 5-7. -23. -7. 29380-5. 23-34. <i>I Sci U S A</i> 90, 5954-8. 03-34. 996) Oncogene 13, 2527 e 15, 1727-36.		

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 nonfat dry milk, 1X TBS, 0.1% Tween® 20 at 4°C with gentle shaking, overnight.		
Applications Key	W: Western Blotting		
Cross-Reactivity Key	H: Human		
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