## #4309 store at -20C

## Phospho-Histone H2A.X (Ser139) (20E3) Rabbit mAb (Biotinylated)



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Applications: W	<b>Reactivity:</b> H M R Mk	<b>Sensitivity:</b> Endogenous	<b>MW (kDa):</b> 15	<b>Source/Isotype:</b> Rabbit IgG	<b>UniProt ID:</b> #P16104	Entrez-Gene Id: 3014	
Product Usage Information		<b>Application</b> Western Blotting			Dilution 1:1000		
Storage		Supplied in 136 mM N 50% glycerol. Store at		2 mM sodium phosphate lot the antibodies.	(pH 7.4) dibasic, 2 r	ng/ml BSA, and	
Specificity/Sensitivity		Phospho-Histone H2A.X (Ser139) (20E3) Rabbit mAb (Biotinylated) detects endogenous levels of H2A.X only when phosphorylated at Ser139.					
Source / Purification		Monoclonal antibody is produced by immunizing animals with a synthetic phosphopeptide corresponding to residues surrounding Ser139 of human H2A.X protein.					
Description		This Cell Signaling Technology antibody is conjugated to biotin under optimal conditions. The biotinylated antibody is expected to exhibit the same species cross-reactivity as the unconjugated antibody (Phospho-Histone H2A.X (Ser139) (20E3) Rabbit mAb #9718).					
Background		in normal human fibro repair following doubl or radiomimetic agent including ATM, ATR, ar at Ser139 at sites of D response is required f MDC1, NBS1, RAD50, I is required for DNA fra response to apoptotic receptor activation, c- response to serum sta cells by WSTF (William concurrent with phosy recruited EYA1 and EYA of DNA repair proteins appears to determine the recruitment of DN Mouse embryonic fibr repair proteins over a	bblasts (1). H2A.X is le-stranded DNA br ts, results in rapid p nd DNA-PK (2,3). Wi NA damage to gen or recruitment of a MRE11, 53BP1, and agmentation during signals. H2A.X is p Jun N-terminal Kina arvation (5-8). H2A.X s-Beuren syndrome oborylation of Ser1 A3 phosphatases (5 s and apoptotic proteins a oblasts expressing poptotic proteins, s he balance of H2A.	presents approximately required for checkpoint eaks (1). DNA damage, c phosphorylation of H2A.) thin minutes following E erate γ-H2A.X (4). This ve multitude of DNA-dama BRCA1 (1). In addition to g apoptosis and is phosp hosphorylated at Ser139 ase (JNK1) in response to X is constitutively phosph e transcription factor) (9, 39, Tyr142 is dephospho 0). While phosphorylation teins to sites of DNA dar ns are recruited. Phosph nd promotes binding of only mutant H2A.X Y142 show a reduced apoptoti X Tyr142 phosphorylation after DNA damage.	-mediated cell cycle aused by ionizing ra ( at Ser139 by PI3K- DNA damage, H2A.X rry early event in the ge response proteir b its role in DNA-dar horylated by variou UV-A irradiation, ar norylated on Tyr142 10). Upon DNA dam rylated at sites of DI n at Ser139 facilitate mage, phosphorylat porylation of H2A.X a pro-apoptotic factor 2F, which favors recr c response to ionizir	arrest and DNA adiation, UV-light, like kinases, is phosphorylated DNA-damage nage repair, H2A.X s kinases in onse to cell death ad p38 MAPK in in undamaged age, and NA damage by s the recruitment ion at Tyr142 at Tyr142 inhibits rs such as JNK1 (9). uitment of DNA ng radiation (9).	
Background Re	ferences	1. Yuan, J. et al. (2010) 2. Rogakou, E.P. et al. 3. Burma, S. et al. (200 4. Rogakou, E.P. et al. 5. Mukherjee, B. et al. 6. Solier, S. et al. (2009 7. Lu, C. et al. (2008) <i>F</i> 9. Cook, P.J. et al. (2009 10. Xiao, A. et al. (2009)	(1998) <i>J Biol Chem 2</i> (1999) <i>J Cell Biol</i> 14 (2006) <i>DNA Repair</i> ) <i>Mol Cell Biol</i> 29, 6 <i>10l Cell</i> 23, 121-32. <i>EBS Lett</i> 582, 2703- 9) <i>Nature</i> 458, 591-	273, 5858-68. 42462-7. 6, 905-16. <i>(Amst)</i> 5, 575-90. 8-82. 8.			

**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.		
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Cross-Reactivity Key	H: Human M: Mouse R: Rat Mk: Monkey		
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