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## Phospho-Rpb1 CTD (Ser2) (E1Z3G) Rabbit



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

<b>Applications:</b> W, W-S, IP, ChIP, ChIP-seq, C&R, C&T	<b>Reactivity:</b> H M R Mk	<b>Sensitivity:</b> Endogenous	<b>MW (kDa):</b> 250	Source/Isotype: Rabbit IgG	<b>UniProt ID:</b> #P24928	Entrez-Gene Id: 5430		
Product Usage Information		For optimal ChIP and ChIP-seq results, use 10 μl of antibody and 10 μg of chromatin (approximately 4 x 10 <sup>6</sup> cells) per IP. This antibody has been validated using SimpleChIP <sup>®</sup> Enzymatic Chromatin IP Kits.						
	The CUT&RUN dilution was determined using CUT&RUN Assay Kit #86652.							
		The CUT&Tag dilution was determined using CUT&Tag Assay Kit #77552.						
		<b>Application</b> Western Blotting			Dilution 1:1000			
		Simple Western™			1:10 - 1:50			
	Immunoprecipitation			1:50				
		Chromatin IP			1:50			
		Chromatin IP-seq			1:50			
		CUT&RUN			1:50			
		CUT&Tag			1:50			
Storage				5), 150 mM NaCl, 100 µg, ot aliquot the antibody.	/ml BSA, 50% glycei	rol and less than		
Specificity/Sen	sitivity	Phospho-Rpb1 CTD (Ser2) (E1Z3G) Rabbit mAb recognizes endogenous levels of Rpb1 only when the carboxy-terminal domain (CTD) heptapeptide repeat [Tyr1, Ser2, Pro3, Thr4, Ser5, Pro6, Ser7] is phosphorylated at Ser2. This antibody does not cross-react with Rpb1 CTD phosphorylated at Ser5 or Ser7.						
Species predict based on 100% homology	ed to react sequence	Hamster, D. melanoga	aster, Xenopus, Zeb	rafish, Bovine, Pig, S. cer	evisiae, C. elegans			
Source / Purific	ation	Monoclonal antibody is produced by immunizing animals with a synthetic phosphopeptide corresponding to residues surrounding Ser2 of the human Rpb1 CTD heptapeptide repeat.						
Background		polymerase, catalyzin as substrates (1). The (POLR2A), contains a u repeated up to 52 tim repeat is subject to m polymerase complex. transcription with chr chromatin modifying transcription initiation through interactions w of RNAPII from gene p transcription factor III enzymes, in addition to initiation and chroma intrinsic pause site, w point, RNAPII is unsta transcription elongati transcription elongati elongation complex a to histone H3 Lys36 m (7,8). Ser2/Ser5-phosp	g the transcription largest subunit, RN unique heptapeptic les in the carboxy-te ultiple post-transla Phosphorylation of omatin remodeling enzymes and RNA n, RNAPII contains a with DNA-bound tra promoters requires H (TFIIH) (2). Phosp to histone H3 Lys4 tin structure (3,4). A here it is halted by ble and frequently on requires phosph on factor P-TEFb (6 nd facilitates recruinethyltransferases, phorylated RNAPII t	Iti-protein complex that of DNA into RNA using t APII subunit B1 (Rpb1), le sequence (Tyr1,Ser2,P erminal domain (CTD) of tional modifications, whi f the CTD during the acti and nascent RNA proce processing proteins to the hypophosphorylated C anscription factors and tl phosphorylation at Ser5 horylation at Ser5 media methyltransferases, whice fithe negative elongation aborts transcription and horylation at Ser2 by CDP ). Phosphorylation at Ser tment of RNA splicing ar which function to promo- hen transcribes the entii II dissociates from the D	he four ribonucleos also known as RNAI ro3,Thr4,Ser5,Pro6, the protein (1). This ch dictate the funct ve transcription cyc ssing by regulating the transcribed gene TD and is recruited the Mediator comple 5 by CDK7, the catal tes the recruitmen the the recruitmen hunction to regul NAPII proceeds dou factors NELF and D dissociates from th (9, the catalytic sub 2 creates a stable t the polyadenylation one length of the ger	side triphosphates PII subunit A Ser7), which is s CTD heptapeptide tional state of the le integrates the recruitment of (1). During to gene promoters ex (1). The escape ytic subunit of t of RNA capping ate transcription wn the gene to an SIF (5). At this he gene. Productive unit of the positive ranscription factors, in addition patible chromatin he to the 3' end,		

Background References	<ul> <li>hypophosphorylated form by various CTD phosphatases (1).In addition to Ser2/Ser5 phosphorylation, Ser7 of the CTD heptapeptide repeat is also phosphorylated during the active transcription cycle. Phosphorylation at Ser7 is required for efficient transcription of small nuclear (sn) RNA genes (9,10). snRNA genes, which are neither spliced nor poly-adenylated, are structurally different from protein-coding genes. Instead of a poly(A) signal found in protein-coding RNAs, snRNAs contain a conserved 3'-box RNA processing element, which is recognized by the Integrator snRNA 3' end processing complex (11,12). Phosphorylation at Ser7 by CDK7 during the early stages of transcription facilitates recruitment of RPAP2, which dephosphorylates Ser5, creating a dual Ser2/Ser7 phosphorylation mark that facilitates recruitment of the Integrator complex and efficient processing of nascent snRNA transcripts (13-15).</li> <li>1. Brookes, E. and Pombo, A. (2009) <i>EMBO Rep</i> 10, 1213-9.</li> <li>2. Komarnitsky, P. et al. (2000) <i>Genes Dev</i> 14, 2452-60.</li> <li>3. Ho, C.K. and Shuman, S. (1999) <i>Mol Cell</i> 3, 405-11.</li> <li>4. Ng, H.H. et al. (2003) <i>Mol Cell</i> 11, 709-19.</li> <li>5. Cheng, B. and Price, D.H. (2007) <i>J Biol Chem</i> 282, 21901-12.</li> <li>6. Marshall, N.F. et al. (1996) <i>J Biol Chem</i> 271, 27176-83.</li> <li>7. Krogan, N.J. et al. (2002) <i>Cell</i> 108, 501-12.</li> <li>9. Chapman, R.D. et al. (2007) <i>Science</i> 318, 1779-9.</li> <li>11. Egloff, S. et al. (2007) <i>Science</i> 318, 1779-9.</li> <li>12. Egloff, S. et al. (2003) <i>Biol Chem</i> 287, 0590-4.</li> <li>13. Akhtar, M.S. et al. (2009) <i>Mol Cell</i> 34, 387-93.</li> <li>14. Egloff, S. et al. (201) <i>J Biol Chem</i> 285, 02564-9.</li> <li>15. Egloff, S. et al. (201) <i>J Biol Chem</i> 285, 02564-9.</li> <li>15. Egloff, S. et al. (2012) <i>Mol Cell</i> 45, 111-22.</li> </ul>		
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	W: Western Blotting W-S: Simple Western™ IP: Immunoprecipitation ChIP: Chromatin IP ChIP-seq: Chromatin IP-seq C&R: CUT&RUN C&T: CUT&Tag		
Cross-Reactivity Key	H: Human M: Mouse R: Rat Mk: Monkey		
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