Revision 8							
Rpb1 CT	D (4H8) N	louse mAb				I Signaling	
Stor					Orders:	877-616-CELL (2355) orders@cellsignal.com	
D					Support:	877-678-TECH (8324)	
#2629					Web:	info@cellsignal.com cellsignal.com	
#				3 Trask Lane D)anvers Massac	husetts 01923 USA	
	ly. Not for Use	in Diagnostic Proced	ures.				
Applications: W, W-S, IP, ChIP, ChIP-seq, C&R, C&T	Reactivity: H M R Mk	Sensitivity: Endogenous	MW (kDa): 250	Source/Isotype: Mouse IgG1	UniProt ID: #P24928	Entrez-Gene Id: 5430	
Product Usage Information							
The CUT&Tag dilution was determined using CUT&Tag Assay Kit #77552.							
		Application Western Blotting Simple Western™ Immunoprecipitation Chromatin IP Chromatin IP-seq CUT&RUN CUT&RUN			Dilution 1:1000 1:50 - 1:250 1:50 1:50 1:50 1:50 1:50		
Storage				5), 150 mM NaCl, 100 µg ot aliquot the antibody		cerol and less than	
Specificity/Sensitivity		Rpb1 CTD (4H8) Antibody detects endogenous levels of total Rpb1 protein (both phosphorylated and unphosphorylated forms).					
Species predicted to react based on 100% sequence homology		Hamster, D. melanogaster, S. cerevisiae					
Source / Purifica	ition			nunizing animals with a 3, Thr4, Ser5, Pro6, Ser7			
Background		RNA polymerase II (RNAPII) is a large multi-protein complex that functions as a DNA-dependent RNA polymerase, catalyzing the transcription of DNA into RNA using the four ribonucleoside triphosphates as substrates (1). The largest subunit, RNAPII subunit B1 (Rpb1), also known as RNAPII subunit A (POLR2A), contains a unique heptapeptide sequence (Tyr1,Ser2,Pro3,Thr4,Ser5,Pro6,Ser7), which is repeated up to 52 times in the carboxy-terminal domain (CTD) of the protein (1). This CTD heptapeptide repeat is subject to multiple post-translational modifications, which dictate the functional state of the polymerase complex. Phosphorylation of the CTD during the active transcription cycle integrates transcription with chromatin remodeling and nascent RNA processing by regulating the recruitment of chromatin modifying enzymes and RNA processing proteins to the transcribed gene (1). During transcription initiation, RNAPII contains a hypophosphorylated CTD and is recruited to gene promoters through interactions with DNA-bound transcription factors and the Mediator complex (1). The escape of RNAPII from gene promoters requires phosphorylation at Ser5 by CDK7, the catalytic subunit of transcription factor IIH (TFIIH) (2). Phosphorylation at Ser5 by CDK7, the catalytic subunit of initiation and chromatin structure (3,4). After promoter escape, RNAPII proceeds down the gene to an intrinsic pause site, where it is halted by the negative elongation factors NELF and DSIF (5). At this point, RNAPII is unstable and frequently aborts transcription and dissociates from the gene. Productive transcription elongation requires phosphorylation at Ser2 by CDK9, the catalytic subunit of the positive transcription elongation factor P-TEFb (6). Phosphorylation at Ser2 creates a stable transcription elongation factor securitement of RNA splicing and polyadenylation factors, in addition to histone H3 Lys36 methyltransferases, which function to promote elongation-compatible chromatin (7,8). Ser2/Ser5-phosphorylated RNAPII then transcribes the enti					

where transcription is terminated. RNAPII dissociates from the DNA and is recycled to the hypophosphorylated form by various CTD phosphatases (1).In addition to Ser2/Ser5 phosphorylation,

Background References	 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). 1. Brookes, E. and Pombo, A. (2009) <i>EMBO Rep</i> 10, 1213-9.
	 Komarnitsky, P. et al. (2000) <i>Genes Dev</i> 14, 2452-60. Ho, C.K. and Shuman, S. (1999) <i>Mol Cell</i> 3, 405-11. Ng, H.H. et al. (2003) <i>Mol Cell</i> 11, 709-19. Cheng, B. and Price, D.H. (2007) <i>J Biol Chem</i> 282, 21901-12. Marshall, N.F. et al. (1996) <i>J Biol Chem</i> 271, 27176-83. Krogan, N.J. et al. (2003) <i>Mol Cell Biol</i> 23, 4207-18. Proudfoot, N.J. et al. (2002) <i>Cell</i> 108, 501-12. Chapman, R.D. et al. (2007) <i>Science</i> 318, 1780-2. Egloff, S. et al. (2003) <i>Biolchem</i> 50, 590-4. Baillat, D. et al. (2005) <i>Cell</i> 123, 265-76. Akhtar, M.S. et al. (2000) <i>Mol Cell</i> 34, 387-93. Egloff, S. et al. (2010) <i>J Biol Chem</i> 285, 20564-9. Egloff, S. et al. (2012) <i>Mol Cell</i> 45, 111-22.
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 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
Trademarks and Patents	Cell Signaling Technology is a trademark of Cell Signaling Technology, Inc.
	All other trademarks are the property of their respective owners. Visit cellsignal.com/trademarks for more information.
Limited Uses	Except as otherwise expressly agreed in a writing signed by a legally authorized representative of CST, the following terms apply to Products provided by CST, its affiliates or its distributors. Any Customer's terms and conditions that are in addition to, or different from, those contained herein, unless separately accepted in writing by a legally authorized representative of CST, are rejected and are of no force or effect.
	Products are labeled with For Research Use Only or a similar labeling statement and have not been approved, cleared, or licensed by the FDA or other regulatory foreign or domestic entity, for any purpose. Customer shall not use any Product for any diagnostic or therapeutic purpose, or otherwise in any manner that conflicts with its labeling statement. Products sold or licensed by CST are provided for Customer as the end-user and solely for research and development uses. Any use of Product for diagnostic, prophylactic or therapeutic purposes, or any purchase of Product for resale (alone or as a component) or other commercial purpose, requires a separate license from CST. Customer shall (a) not sell, license, loan, donate or otherwise transfer or make available any Product to any third party, whether alone or in combination with other materials, or use the Products to manufacture any commercial products, (b) not copy, modify, reverse engineer, decompile, disassemble or otherwise attempt to discover the underlying structure or technology of the Products, or use the Products for the purpose of developing any products or services that would compete with CST products or services, (c) not alter or remove from the Products solely in accordance with CST Product Terms of Sale and any applicable documentation, and (e) comply with any license, terms of service or similar agreement with respect to any third party products or services used by Customer in connection with the Products.