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Phospho-PKCδ (Ser359) (D2X1P) Rabbit mAb



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Applications: W	Reactivity: H M	Sensitivity: Endogenous	MW (kDa): 78	Source/Isotype: Rabbit IgG	UniProt ID: #Q05655	Entrez-Gene Id: 5580	
Product Usage Information		Application Western Blotting			Dilution 1:1000		
Storage				i), 150 mM NaCl, 100 μg/ ot aliquot the antibody.	/ml BSA, 50% glycer	ol and less than	
Specificity/Sensitivity		Phospho-PKCδ (Ser359) (D2X1P) Rabbit mAb recognizes endogenous levels of PKCδ protein only when phosphorylated at Ser359.					
Species predicto based on 100% homology	ed to react sequence	Rat					
Source / Purific	ation	Monoclonal antibody is produced by immunizing animals with a synthetic phosphopeptide corresponding to residues surrounding Ser359 of human PKC δ protein.		eptide			
Background		Activation of protein kinase C (PKC) is one of the earliest events in a cascade that controls a variety of cellular responses, including secretion, gene expression, proliferation, and muscle contraction (1,2). PKC isoforms belong to three groups based on calcium dependency and activators. Classical PKCs are calcium-dependent via their C2 domains and are activated by phosphatidylserine (PS), diacylglycerol (DAG), and phorbol esters (TPA, PMA) through their cysteine-rich C1 domains. Both novel and atypical PKCs are calcium-independent, but only novel PKCs are activated by PS, DAG, and phorbol esters (3-5). Members of these three PKC groups contain a pseudo-substrate or autoinhibitory domain that binds to substrate-binding sites in the catalytic domain to prevent activation in the absence of cofactors or activators. Control of PKC activity is regulated through three distinct phosphorylation events. Phosphorylation occurs <i>in vivo</i> at Thr500 in the activation loop, at Thr641 through autophosphorylation, and at the carboxy-terminal hydrophobic site Ser660 (2). Atypical PKC isoforms lack hydrophobic region phosphorylation, which correlates with the presence of glutamic acid rather than the serine or threonine residues found in more typical PKC superfamily is PKCμ (PKD), which is regulated by DAG and TPA through its C1 domain. PKD is distinguished by the presence of a PH domain and by its unique substrate recognition and Golgi localization (6). PKC-related kinases (PRK) lack the C1 domain and do not respond to DAG or phorbol esters. Phosphatidylinositol lipids activate PRKs, and small Rho-family GTPases bind to the homology region 1 (HR1) to regulate PRK kinase activity (7).					
Background Re	ferences	1. Nishizuka, Y. (1984) 2. Keranen, L.M. et al. 3. Mellor, H. and Parke 4. Ron, D. and Kazanie 5. Moscat, J. and Diaz- 6. Baron, C.L. and Mal 7. Flynn, P. et al. (2000	(1995) <i>Curr Biol</i> 5, er, P.J. (1998) <i>Bioche</i> etz, M.G. (1999) <i>FASI</i> Meco, M.T. (2000) <i>E</i> hotra, V. (2002) <i>Scie</i>	m J 332 (Pt 2), 281-92. EB J 13, 1658-76. MBO Rep 1, 399-403. ence 295, 325-8.			
Species Reactiv	ity	Species reactivity is de	etermined by testin	g in at least one approve	ed application (e.g.,	western blot).	
Western Blot Bu	uffer			membrane with diluted with gentle shaking, ove		ר 5% w/v nonfat	
Applications Ke	у	W: Western Blotting					
Cross-Reactivity	у Кеу	H: Human M: Mouse					

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