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## CdGAP (D6J9G) Rabbit mAb



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Applications: W	<b>Reactivity:</b> H M R Mk	<b>Sensitivity:</b> Endogenous	<b>MW (kDa):</b> 250	<b>Source/Isotype:</b> Rabbit IgG	UniProt ID: #Q2M1Z3	Entrez-Gene Id: 57514		
Product Usage Information	9	Application Dilution   Western Blotting 1:1000						
Storage		Supplied in 10 mM sodium HEPES (pH 7.5), 150 mM NaCl, 100 μg/ml BSA, 50% glycerol and less than 0.02% sodium azide. Store at –20°C. Do not aliquot the antibody.						
Specificity/Sensitivity		CdGAP (D6J9G) Rabbit mAb recognizes endogenous levels of total CdGAP protein.						
Source / Purification		Monoclonal antibody is produced by immunizing animals with a synthetic peptide corresponding to residues surrounding Ser520 of human CdGAP protein.						
Background		The Rho family of small GTPases, including Rho, Rac, and Cdc42, act as molecular switches that regulate processes such as cell migration, adhesion, proliferation, and differentiation. They are activated by guanine nucleotide exchange factors (GEFs), which catalyze the exchange of bound GDP for GTP, and inhibited by GTPase activating proteins (GAPs), which catalyze the hydrolysis of GTP to GDP (1). The serine- and proline-rich GAP protein, Cdc42 GAP (CdGAP), has been shown to be a negative regulator of both Cdc42 and Rac1, but not RhoA (2,3). This protein contains three domains: an amino-terminal GAP domain, a central domain, and a carboxy-terminal proline-rich domain containing five Src homology 3 (SH3)-binding sites. It is suggested that threonine and serine phosphorylation within the proline-rich domain likely alters protein-protein interactions and determines the localization of CdGAP (4). Phosphorylation of CdGAP on threonine 776 by both ERK-1 and GSK-3 has been shown to negatively regulate protein activity, possibly by inducing a conformational change within the protein disrupting its ability to bind SH3 domains (4,5). Upregulation of CdGAP has been shown to increase cell proliferation and it has been suggested that this protein may play a role in TGF-β-induced cell growth, motility, and invasion in some breast cancer cells (6).						
Background Ro	eferences	1. Takai, Y. et al. (2001) <i>Physiol Rev</i> 81, 153-208. 2. Tcherkezian, J. et al. (2006) <i>Biol Cell</i> 98, 445-56. 3. Lamarche-Vane, N. and Hall, A. (1998) <i>J Biol Chem</i> 273, 29172-7. 4. Tcherkezian, J. et al. (2005) <i>Mol Cell Biol</i> 25, 6314-29. 5. Danek, E.I. et al. (2007) <i>J Biol Chem</i> 282, 3624-31. 6. He, Y. et al. (2011) <i>Oncogene</i> 30, 1032-45.						
Species Reacti	vitv	Species reactivity is de	etermined by testing	g in at least one approve	ed application (e.g.,	western blot).		
Western Blot E	-	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 K	ey	W: Western Blotting						
Cross-Reactivit	ty Key	H: Human M: Mouse R: Rat Mk: Monkey						
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