Phospho-ALK (Tyr1586) Antibody





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

Applications: W	Reactivity: H	Sensitivity: Endogenous	MW (kDa): 80 (NPM-ALK) 220 (ALK)	Source/Isotype: Rabbit	UniProt ID: #Q9UM73	Entrez-Gene Id: 238		
Product Usage Information	9	Application Western Blotting			Dilution 1:1000			
Storage	StorageSupplied in 10 mM sodium HEPES (pH 7.5), 20°C. Do not aliquot the antibody.), 150 mM NaCl, 100 $\mu g/ml$ BSA and 50% glycerol. Store at –			
Specificity/Sensitivity		Phospho-ALK (Tyr1586) Antibody detects ALK only when phosphorylated at tyrosine 1586 (equivalent to Tyr646 of NPM-ALK).						
Source / Purifi	cation	Polyclonal antibodies are produced by immunizing animals with a synthetic phosphopeptide corresponding to residues surrounding Tyr1586 of human ALK. Antibodies are purified by protein A and peptide affinity chromatography.						
Background	 Background Anaplastic lymphoma kinase (ALK) is a tyrosine kinase receptor for pleiotrophin (PTN), a growth involved in embryonic brain development (1-3). In ALK-expressing cells, PTN induces phosphory of both ALK and the downstream effectors IRS-1, Shc, PLCy, and PI3 kinase (1). ALK was originall discovered as a nucleophosmin (NPM)-ALK fusion protein produced by a translocation (4). Invest have found that the NPM-ALK fusion protein is a constitutively active, oncogenic tyrosine kinase associated with anaplastic lymphoma (4). Research literature suggests that activation of PLCy by ALK may be a crucial step for its mitogenic activity and involved in the pathogenesis of anaplast lymphomas (5). A distinct ALK oncogenic fusion protein involving ALK and echinoderm microtubule-associated plike 4 (EML4) has been described in the research literature from a non-small cell lung cancer (NS cell line, with corresponding fusion transcripts present in some cases of lung adenocarcinoma. short, amino-terminal region of the microtubule-associated protein EML4 is fused to the kinase of ALK (6-8). 					phosphorylation vas originally on (4). Investigators osine kinase n of PLCγ by NPM- of anaplastic associated protein g cancer (NSCLC) carcinoma. The		
Background R	eferences	 Stoica, G.E. et al. (2001) <i>J Biol Chem</i> 276, 16772-9. Iwahara, T. et al. (1997) <i>Oncogene</i> 14, 439-49. Morris, S.W. et al. (1997) <i>Oncogene</i> 14, 2175-88. Morris, S.W. et al. (1994) <i>Science</i> 263, 1281-4. Bai, R.Y. et al. (1998) <i>Mol Cell Biol</i> 18, 6951-61. Rikova, K. et al. (2007) <i>Cell</i> 131, 1190-203. Takeuchi, K. et al. (2008) <i>Clin Cancer Res</i> 14, 6618-24. Soda, M. et al. (2007) Nature 448, 561-6. 						
Species Reacti	vity	Species reactivity is	determined by testing	in at least one approve	ed application (e.g.,	western blot).		
Western Blot E	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.				n 5% w/v BSA, 1X		
Applications K	ey	W: Western Blotting						
Cross-Reactivi	ty Key	H: Human						
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