Cbl (phospho Tyr700) Polyclonal Antibody

Catalog No: #13998

Package Size: #13998-1 50ul #13998-2 100ul



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Product Name	Cbl (phospho Tyr700) Polyclonal Antibody	
Host Species	Rabbit	
Purification	The antibody was affinity-purified from rabbit antiserum by affinity-chromatography using epitope-specific	
	immunogen.	
Applications	WB,IHC-p,IF/ICC,ELISA	
Species Reactivity	Human,Mouse,Rat	
Specificity	Phospho-Cbl (Y700) Polyclonal Antibody detects endogenous levels of Cbl protein only when phosphorylated	
	at Y700.	
Immunogen Description	The antiserum was produced against synthesized peptide derived from human CBL around the	
	phosphorylation site of Tyr700. AA range:666-715	
Other Names	CBL; CBL2; RNF55; E3 ubiquitin-protein ligase CBL; Casitas B-lineage lymphoma proto-oncogene;	
	Proto-oncogene c-Cbl; RING finger protein 55; Signal transduction protein CBL	
Accession No.	Swiss Prot:P22681GeneID:867	
SDS-PAGE MW	120	
Concentration	1 mg/ml	
Formulation	Liquid in PBS containing 50% glycerol, 0.5% BSA and 0.02% sodium azide.	
Storage	-20°C/1	

Application Details

Western Blot: 1/500 - 1/2000. Immunohistochemistry: 1/100 - 1/300. Immunofluorescence: 1/200 - 1/1000. ELISA: 1/10000. Not yet tested in other applications.

Background

Cbl proto-oncogene(CBL) Homo sapiens This gene is a proto-oncogene that encodes a RING finger E3 ubiquitin ligase. The encoded protein is one of the enzymes required for targeting substrates for degradation by the proteasome. This protein mediates the transfer of ubiquitin from ubiquitin conjugating enzymes (E2) to specific substrates. This protein also contains an N-terminal phosphotyrosine binding domain that allows it to interact with numerous tyrosine-phosphorylated substrates and target them for proteasome degradation. As such it functions as a negative regulator of many signal transduction pathways. This gene has been found to be mutated or translocated in many cancers including acute myeloid leukaemia, and expansion of CGG repeats in the 5' UTR has been associated with Jacobsen syndrome. Mutations in this gene are also the cause of Noonan syndrome-like disorder. [provided by RefSeq, Jul 2016],

Note: This product is for in vitro research use only and is not intended for use in humans or animals.