

CD42b rabbit pAb

Cat No.:ES4123

For research use only

Overview

Product Name	CD42b rabbit pAb
Host species	Rabbit
Applications	WB;IHC;IF;ELISA
Species Cross-Reactivity	Human;Rat;Mouse;
Recommended dilutions	Western Blot: 1/500 - 1/2000. IHC-p: 1/100-1/300.
	ELISA: 1/20000. Not yet tested in other applications.
Immunogen	The antiserum was produced against synthesized
	peptide derived from the Internal region of human
	GP1BA. AA range:271-320
Specificity	CD42b Polyclonal Antibody detects endogenous
	levels of CD42b protein.
Formulation	Liquid in PBS containing 50% glycerol, 0.5% BSA and
	0.02% sodium azide.
Storage	Store at -20°C. Avoid repeated freeze-thaw cycles.
Protein Name	Platelet glycoprotein Ib alpha chain
Gene Name	GP1BA
Cellular localization	Membrane; Single-pass type I membrane protein.
Purification	The antibody was affinity-purified from rabbit
	antiserum by affinity-chromatography using
	epitope-specific immunogen.
Clonality	Polyclonal
Concentration	1 mg/ml
Observed band	69kD
Human Gene ID	2811
Human Swiss-Prot Number	P07359
Alternative Names	GP1BA; Platelet glycoprotein Ib alpha chain; GP-Ib
	alpha; GPIb-alpha; GPIbA; Glycoprotein Ibalpha;
	Antigen CD42b-alpha; CD42b
Background	Glycoprotein Ib (GP Ib) is a platelet surface
	membrane glycoprotein composed of a
	heterodimer, an alpha chain and a beta chain, that is
	linked by disulfide bonds. The Gp Ib functions as a
	receptor for von Willebrand factor (VWF). The



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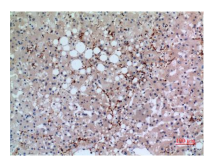
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complete receptor complex includes noncovalent association of the alpha and beta subunits with platelet glycoprotein IX and platelet glycoprotein V. The binding of the GP Ib-IX-V complex to VWF facilitates initial platelet adhesion to vascular subendothelium after vascular injury, and also initiates signaling events within the platelet that lead to enhanced platelet activation, thrombosis, and hemostasis. This gene encodes the alpha subunit. Mutations in this gene result in Bernard-Soulier syndromes and platelet-type von Willebrand disease. The coding region of this gene is known to contain a polymophic variable number tandem repeat (VNTR) domain that is

Western Blot analysis of HeLa cells using CD42b Polyclonal Antibody. Secondary antibody(catalog#:RS0002) was diluted at 1:20000

Immunohistochemical analysis of paraffin-embedded human-liver, antibody was diluted at 1:100



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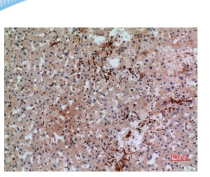
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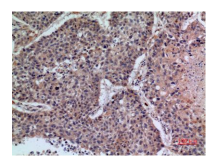
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Immunohistochemical analysis of paraffin-embedded human-liver, antibody was diluted at 1:100



Immunohistochemical analysis of paraffin-embedded human-lung, antibody was diluted at 1:100





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