



# SPRE1 Polyclonal Antibody

<b>Catalog No</b>	BYab-07258
<b>Isotype</b>	IgG
<b>Reactivity</b>	Human;Mouse
<b>Applications</b>	WB;ELISA
<b>Gene Name</b>	SPRED1
<b>Protein Name</b>	Sprouty-related, EVH1 domain-containing protein 1 (Spred-1) (hSpred1)
<b>Immunogen</b>	Synthesized peptide derived from human protein . at AA range: 210-290
<b>Specificity</b>	SPRE1 Polyclonal Antibody detects endogenous levels of protein.
<b>Formulation</b>	Liquid in PBS containing 50% glycerol, and 0.02% sodium azide.
<b>Source</b>	Polyclonal, Rabbit,IgG
<b>Purification</b>	The antibody was affinity-purified from rabbit antiserum by affinity-chromatography using epitope-specific immunogen.
<b>Dilution</b>	WB 1:500-2000 ELISA 1:5000-20000
<b>Concentration</b>	1 mg/ml
<b>Purity</b>	≥90%
<b>Storage Stability</b>	-20°C/1 year
<b>Synonyms</b>	
<b>Observed Band</b>	48kD
<b>Cell Pathway</b>	Cell membrane ; Peripheral membrane protein . Membrane, caveola ; Peripheral membrane protein . Nucleus . Localized in cholesterol-rich membrane raft/caveola fractions.
<b>Tissue Specificity</b>	Weakly expressed in embryonic cell line HEK293.
<b>Function</b>	disease:Defects in SPRED1 are the cause of neurofibromatosis type 1-like syndrome (NFLS) [MIM:611431]. Neurofibromatosis type 1 (NF1) is one of the most frequent autosomal dominant diseases. It belongs to the group of disorders known as the 'neuro-cardio-facial-cutaneous' syndromes, present with a variable degree of cognitive impairment, facial dysmorphism, congenital heart defects and skin abnormalities. NFLS is a form of these disorders with autosomal dominant trait consisting of multiple cafe-au-lait spots, axillary freckling, macrocephaly and a Noonan-like dysmorphism in some individuals.,function:Tyrosine kinase substrate that inhibits growth-factor-mediated activation of MAP kinase. Negatively regulates hematopoiesis of bone marrow.,PTM:Phosphorylated on tyrosine.,sequence caution:Contaminating sequence. Potential poly-A

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sequence.,similarity:Contains 1 KBD domain.,similarity:Contains

**Background**

The protein encoded by this gene is a member of the Sprouty family of proteins and is phosphorylated by tyrosine kinase in response to several growth factors. The encoded protein can act as a homodimer or as a heterodimer with SPRED2 to regulate activation of the MAP kinase cascade. Defects in this gene are a cause of neurofibromatosis type 1-like syndrome (NFLS). [provided by RefSeq, Jul 2008],

**matters needing attention**

Avoid repeated freezing and thawing!

**Usage suggestions**

This product can be used in immunological reaction related experiments. For more information, please consult technical personnel.

**Products Images**