

Filamin A (PT0388R) PT® Rabbit mAb

Catalog No :	YM8238
Reactivity :	Human; Mouse; Rat;
Applications :	WB;IHC;IF;IP;ELISA
Target :	Filamin 1
Fields :	>>MAPK signaling pathway;>>Focal adhesion;>>Salmonella infection;>>Proteoglycans in cancer
Gene Name :	FLNA
Protein Name :	Filamin-A
Human Gene Id :	2316
Human Swiss Prot	P21333
Mouse Gene Id :	192176
Mouse Swiss Prot	Q8BTM8
No : Specificity :	endogenous
Formulation :	PBS, 50% glycerol, 0.05% Proclin 300, 0.05%BSA
Source :	Monoclonal, rabbit, IgG, Kappa
Dilution :	IHC 1:4000-1:10000,WB 1:1000-1:5000,IF 1:200-1:1000,ELISA 1:5000-1:20000,IP 1:50-1:200,
Purification :	Protein A
Storage Stability :	-15°C to -25°C/1 year(Do not lower than -25°C)
Molecularweight :	281kD



Best Tools for immunology Research

Observed Band : 281kD

Cell Pathway : MAPK_ERK_Growth;MAPK_G_Protein;Focal adhesion;

Background : filamin A(FLNA) Homo sapiens The protein encoded by this gene is an actinbinding protein that crosslinks actin filaments and links actin filaments to membrane glycoproteins. The encoded protein is involved in remodeling the cytoskeleton to effect changes in cell shape and migration. This protein interacts with integrins, transmembrane receptor complexes, and second messengers. Defects in this gene are a cause of several syndromes, including periventricular nodular heterotopias (PVNH1, PVNH4), otopalatodigital syndromes (OPD1, OPD2), frontometaphyseal dysplasia (FMD), Melnick-Needles syndrome (MNS), and X-linked congenital idiopathic intestinal pseudoobstruction (CIIPX). Two transcript variants encoding different isoforms have been found for this gene.[provided by RefSeq, Mar 2009],

Function :

disease:Defects in FLNA are associated with cerebrofrontofacial syndrome [MIM:608578]. This syndrome consists of a phenotype of male PVNH, with relatively normal development, no epilepsy or other neurological abnormality, severe constipation, and facial dysmorphism and without a discernible skeletal phenotype.,disease:Defects in FLNA are the cause of frontometaphyseal dysplasia (FMD) [MIM:305620]. FMD is a congenital bone disease characterized by supraorbital hyperostosis, deafness and digital anomalies.,disease:Defects in FLNA are the cause of Melnick-Needles syndrome (MNS) [MIM:309350]. MNS is a severe congenital bone disorder characterized by typical facies (exophthalmos, full cheeks, micrognathia and malalignment of teeth), flaring of the metaphyses of long bones, s-like curvature of bones of legs, irregular constrictions in the ribs, and sclerosis of base of skull.,disease:Defects i

Subcellular	Cytoplasm
Location :	
Expression :	Ubiquitous.
Tag :	hot,recombinant
Sort :	6065
No4 :	1
Host :	Rabbit
Modifications :	Unmodified

Products Images





Various whole cell lysates were separated by 4-8% SDS-PAGE, and the membrane was blotted with anti-Filamin A (PT0388R) antibody. The HRP-conjugated Goat anti-Rabbit IgG(H + L) antibody was used to detect the antibody. Lane 1: HepG2 Lane 2: Mouse liver Predicted band size: 281kDa Observed band size: 281kDa