

CD61 (ABT134R) rabbit mAb

YM7255 Catalog No:

Reactivity: Human;

IHC; WB; ELISA **Applications:**

Target: Integrin β3

>>Rap1 signaling pathway;>>Phagosome;>>PI3K-Akt signaling Fields:

pathway:>>Osteoclast differentiation:>>Focal adhesion:>>ECM-receptor

interaction;>>Platelet activation;>>Neutrophil extracellular trap formation;>>Hematopoietic cell lineage;>>Regulation of actin

cytoskeleton;>>Thyroid hormone signaling pathway;>>Human cytomegalovirus

infection;>>Human papillomavirus infection;>>Herpes simplex virus 1

Synthesized peptide derived from human CD61 AA range:1-100

infection;>>Proteoglycans in cancer;>>MicroRNAs in cancer;>>Hypertrophic cardiomyopathy;>>Arrhythmogenic right ventricular cardiomyopathy;>>Dilated

cardiomyopathy;>>Fluid shear stress and atherosclerosis

Gene Name: ITGB3

Protein Name: Integrin beta-3 (Platelet membrane glycoprotein IIIa) (GPIIIa) (CD antigen

CD61)

Human Gene Id: 3690

Human Swiss Prot

Immunogen:

No:

P05106

Specificity: This antibody detects endogenous levels of Integrin β3

Formulation: PBS, 50% glycerol, 0.05% Proclin 300, 0.05% BSA

Source: Monoclonal, Rabbit IgG1, Kappa

IHC 1:100-500, WB 1:500-1000, ELISA 1:5000-20000 **Dilution:**

Purification: Recombinant Expression and Affinity purified

1/3



Storage Stability: -15°C to -25°C/1 year(Do not lower than -25°C)

Molecularweight: 87kD

Cell Pathway: Focal adhesion; ECM-receptor interaction; Hematopoietic cell lineage; Regulates

Actin and Cytoskeleton; Hypertrophic cardiomyopathy (HCM); Arrhythmogenic

right ventricular cardiomyopathy (ARVC); Dilated car

Background: The ITGB3 protein product is the integrin beta chain beta 3. Integrins are

integral cell-surface proteins composed of an alpha chain and a beta chain. A given chain may combine with multiple partners resulting in different integrins. Integrin beta 3 is found along with the alpha IIb chain in platelets. Integrins are known to participate in cell adhesion as well as cell-surface mediated signalling.

[provided by RefSeq, Jul 2008],

Function: disease:Defects in ITGB3 are a cause of Glanzmann thrombasthenia (GT)

[MIM:273800]; also known as thrombasthenia of Glanzmann and Naegeli. GT is the most common inherited disease of platelets. Its inheritance is autosomal recessive. It is characterized by mucocutaneous bleeding of mild-to-moderate severity and the inability of this integrin to recognize macromolecular or synthetic peptide ligands. GT has been classified clinically into types I and II. In type I, platelets show absence of the glycoprotein IIb-IIIa complexes at their surface and lack fibrinogen and clot retraction capability. In type II, the platelets express the GPIIb-IIIa complex at reduced levels (5-20% controls), have detectable amounts of fibrinogen, and have low or moderate clot retraction capability. The platelets of GT variants have normal or near normal (60-100%) expression of dysfunctional

receptors.,function:Int

Subcellular Location:

Cytoplasmic

Expression: Isoform beta-3A and isoform beta-3C are widely expressed. Isoform beta-3A is

specifically expressed in osteoblast cells; isoform beta-3C is specifically

expressed in prostate and testis.

Tag: recombinant

Sort: 800

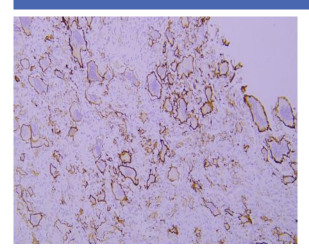
No4: __1

Host: Rabbit

Modifications : Unmodified



Products Images



Immunohistochemical analysis of paraffin-embedded human Giant cell tumor of bone. 1, Antibody was incubated at 4° overnight. 2, Citrate buffer of pH6.0 was used for antigen retrieval. 3,Secondary antibody was diluted at 1:200(room temperature, 30min).