

Flt-4 Monoclonal Antibody

Catalog No: YM0279

Reactivity: Human

Applications: WB;ELISA

Target: VEGFR3

Fields: >>MAPK signaling pathway;>>Ras signaling pathway;>>Rap1 signaling

pathway;>>Calcium signaling pathway;>>PI3K-Akt signaling pathway;>>Focal

adhesion;>>Pathways in cancer;>>Breast cancer

Gene Name: FLT4

Protein Name: Vascular endothelial growth factor receptor 3

P35917

Human Gene Id: 2324

Human Swiss Prot P35916

No:

Mouse Swiss Prot

No:

Immunogen: Purified recombinant fragment of human Flt-4 expressed in E. Coli.

Specificity: Flt-4 Monoclonal Antibody detects endogenous levels of Flt-4 protein.

Formulation : Liquid in PBS containing 50% glycerol, 0.5% BSA and 0.02% sodium azide.

Source: Monoclonal, Mouse

Dilution: WB 1:500 - 1:2000. ELISA: 1:10000. Not yet tested in other applications.

Purification : Affinity purification

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

Molecularweight: 153kD

1/3



Cell Pathway: Cytokine-cytokine receptor interaction; Focal adhesion;

P References : 1. Prostate. 2009 Jun 15;69(9):982-90.

2. J Cell Sci. 2009 Sep 15;122(Pt 18):3358-64.

3. Oncol Rep. 2009 Nov;22(5):1093-100.

Background: This gene encodes a tyrosine kinase receptor for vascular endothelial growth

factors C and D. The protein is thought to be involved in lymphangiogenesis and

maintenance of the lymphatic endothelium. Mutations in this gene cause

hereditary lymphedema type IA. [provided by RefSeq, Jul 2008],

Function : catalytic activity:ATP + a [protein]-L-tyrosine = ADP + a [protein]-L-tyrosine

phosphate., disease: Defects in FLT4 are found in juvenile hemangioma. Juvenile hemangiomas are the most common tumors of infancy, occurring as many as 10% of all births. These benign vascular lesions enlarge rapidly during the first year of life by hyperplasia of endothelial cells and attendant pericytes, and then

spontaneously involute over a period of years, leaving loose fibrofatty

tissue., disease: Defects in FLT4 are the cause of lymphedema hereditary type 1 (LYH1A) [MIM:153100]; also known as Nonne-Milroy lymphedema or Milroy disease. Hereditary lymphedema is a chronic disabling condition which results in

swelling of the extremities due to altered lymphatic flow. Patients with lymphedema suffer from recurrent local infections and physical

impairment.,function:Receptor for VEGFC. Has a tyrosine-protein kinas

Subcellular Location:

Cell membrane ; Single-pass type I membrane protein. Cytoplasm . Nucleus . Ligand-mediated autophosphorylation leads to rapid internalization. .; [Isoform 1]:

Cell membrane; Single-pass type I membrane protein. Ligand-mediated autophosphorylation leads to rapid internalization.; [Isoform 2]: Cell membrane;

Single-pass type I membrane protein.; [Isoform 3]: Secreted. Cytoplasm.

Expression: Detected in endothelial cells (at protein level). Widely expressed. Detected in

fetal spleen, lung and brain. Detected in adult liver, muscle, thymus, placenta,

lung, testis, ovary, prostate, heart, and kidney.

Tag: hot

Sort : 6195

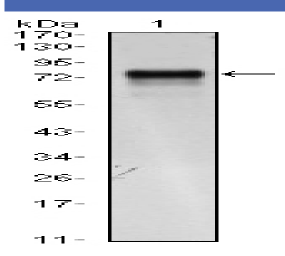
No4: 1

Host: Mouse

Modifications : Unmodified



Products Images



Western Blot analysis using Flt-4 Monoclonal Antibody against FLT4-hlgGFc transfected HEK293 cell lysate.