

FGF Receptor 2 (Phospho Tyr769) rabbit pAb

Catalog No: YP1701

Reactivity: Human; Mouse; Rat

Applications: WB

Target: FGF Receptor 2

Fields: >>EGFR tyrosine kinase inhibitor resistance;>>MAPK signaling pathway;>>Ras

signaling pathway;>>Rap1 signaling pathway;>>Calcium signaling

pathway;>>Endocytosis;>>PI3K-Akt signaling pathway;>>Signaling pathways

regulating pluripotency of stem cells;>>Regulation of actin

cytoskeleton;>>Pathways in cancer;>>Prostate cancer;>>Gastric

cancer;>>Central carbon metabolism in cancer

Gene Name: FGFR2 BEK KGFR KSAM

Protein Name : FGFR2 (Phospho-Tyr769)

P21802

P21803

Human Gene ld: 2263

Human Swiss Prot

No:

Mouse Gene Id: 14183

Mouse Swiss Prot

No:

Immunogen: Synthesized peptide derived from human FGFR2 (Phospho-Tyr769)

Specificity: This antibody detects endogenous levels of FGFR2 (Phospho-Tyr769) at

Human, Mouse, Rat

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

Source: Polyclonal, Rabbit, IgG

Dilution : WB 1:500-2000

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Purification: The antibody was affinity-purified from rabbit serum by affinity-chromatography

using specific immunogen.

Concentration: 1 mg/ml

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

Molecularweight: 90kD

Background: The protein encoded by this gene is a member of the fibroblast growth factor

receptor family, where amino acid sequence is highly conserved between members and throughout evolution. FGFR family members differ from one another in their ligand affinities and tissue distribution. A full-length representative protein consists of an extracellular region, composed of three immunoglobulin-like domains, a single hydrophobic membrane-spanning segment and a cytoplasmic tyrosine kinase domain. The extracellular portion of the protein interacts with fibroblast growth factors, setting in motion a cascade of downstream signals, ultimately influencing mitogenesis and differentiation. This particular family member is a high-affinity receptor for acidic, basic and/or keratinocyte growth factor, depending on the isoform. Mutations in this gene are associated with

Crouzon syndrome, Pfeiffer syndrome, C

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

phosphate., disease: Defects in FGFR2 are a cause of Apert syndrome (APRS) [MIM:101200]; also known as acrocephalosyndactyly type 1 (ACS1). APRS is a syndrome characterized by facio-cranio-synostosis, osseous and membranous syndactyly of the four extremities, and midface hypoplasia. The craniosynostosis is bicoronal and results in acrocephaly of brachysphenocephalic type. Syndactyly of the fingers and toes may be total (mitten hands and sock feet) or partial affecting the second, third, and fourth digits. Intellectual deficit is frequent and

often severe, usually being associated with cerebral

malformations., disease: Defects in FGFR2 are a cause of Jackson-Weiss syndrome (JWS) [MIM:123150]. JWS is an autosomal dominant craniosynostosis syndrome characterized by craniofacial abnormalities and abnormality of the fe

Subcellular Location:

Cell membrane; Single-pass type I membrane protein. Golgi apparatus. Cytoplasmic vesicle. Detected on osteoblast plasma membrane lipid rafts. After ligand binding, the activated receptor is rapidly internalized and degraded.; [Isoform 1]: Cell membrane; Single-pass type I membrane protein. After ligand binding, the activated receptor is rapidly internalized and degraded.; [Isoform 3]:

Cell membrane; Single-pass type I membrane protein. After ligand binding, the activated receptor is rapidly internalized and degraded.; [Isoform 8]: Secreted.;

[Isoform 13]: Secreted.

Expression: Blood, Brain, Cerebellum, Cornea, Mammary gland, Neonatal brain stem, Pla

Sort : 25181



Host: Rabbit	
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Modifications : Phospho

Products Images

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