

Tuberin/TSC2 (Phospho Ser1254) rabbit pAb

Catalog No: YP1538

Reactivity: Human; Mouse; Rat

Applications: WB

Target: Tuberin

Fields: >>Phospholipase D signaling pathway;>>p53 signaling pathway;>>Autophagy -

animal;>>mTOR signaling pathway;>>PI3K-Akt signaling pathway;>>AMPK

signaling pathway;>>Longevity regulating pathway;>>Cellular

senescence;>>Thermogenesis;>>Insulin signaling pathway;>>Thyroid hormone signaling pathway;>>Human cytomegalovirus infection;>>Human papillomavirus infection;>>Herpes simplex virus 1 infection;>>Choline metabolism in cancer

Gene Name: TSC2 TSC4

Protein Name: Tuberin/TSC2 (Ser1254)

Q61037

Human Gene Id: 7249

Human Swiss Prot P49815

No:

Mouse Swiss Prot

No:

Rat Gene Id: 24855

Rat Swiss Prot No: P49816

Immunogen: Synthesized phosho peptide around human Tuberin(Ser1254)

Specificity: This antibody detects endogenous levels of Human Mouse Rat Tuberin/TSC2

(phospho-Ser1254)

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

Source : Polyclonal, Rabbit, IgG

1/3



Dilution: WB 1:1000-2000

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)

Observed Band: 200kD

Cell Pathway: Insulin Receptor; mTOR; B Cell Receptor; PI3K/Akt; AMPK

Background: Mutations in this gene lead to tuberous sclerosis complex. Its gene product is

believed to be a tumor suppressor and is able to stimulate specific GTPases. The protein associates with hamartin in a cytosolic complex, possibly acting as a chaperone for hamartin. Alternative splicing results in multiple transcript variants

encoding different isoforms. [provided by RefSeq, Jul 2008],

Function: alternative products:Additional isoforms seem to exist. Experimental

confirmation may be lacking for some isoforms, disease: Defects in TSC2 are a cause of lymphangioleiomyomatosis (LAM) [MIM:606690]. LAM is a progressive and often fatal lung disease characterized by a diffuse proliferation of abnormal smooth muscle cells in the lungs. It affects almost exclusively young women and can occur as an isolated disorder or in association with tuberous sclerosis

complex.,disease:Defects in TSC2 are the cause of tuberous sclerosis complex (TSC) [MIM:191100]. The molecular basis of TSC is a functional impairment of the tuberin-hamartin complex. TSC is an autosomal dominant multi-system disorder that affects especially the brain, kidneys, heart, and skin. TSC is characterized by hamartomas (benign overgrowths predominantly of a cell or

tissue type that occurs normally in the organ) and hamartias (de

Subcellular Location:

Cytoplasm. Membrane; Peripheral membrane protein. At steady state found in

association with membranes.

Expression: Liver, brain, heart, lymphocytes, fibroblasts, biliary epithelium, pancreas,

skeletal muscle, kidney, lung and placenta.

Sort: 23726

No4: 1

Host: Rabbit

Modifications: Phospho



Products Images