

Met (Phospho Tyr1234/1235) (PT0745R) PT® Rabbit mAb

CatalogNo: YM8554 Recombinant 💦

Key Features

Host Species

Rabbit

MW • 155kD (Calculated) 155kD (Observed) Reactivity

Human,Mouse,Rat

Isotype • IgG,Kappa Applications

WB,IHC,IF,IP,ELISA

Recommended Dilution Ratios

IHC 1:200-1:1000 WB 1:2000-1:10000 IF 1:200-1:1000 ELISA 1:5000-1:20000 IP 1:50-1:200

Storage

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

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

Basic Information

Clonality Monoclonal

Clone Number PT0745R

Immunogen Information

Specificity This antibody detects endogenous levels of Met only when phosphorylated at Tyr1234 or Thr1235,and dually phosphorylated at two sites.

Target Information

Gene name	MET		
Protein Name	Met (Tyr1234/1235) Organism	Gene ID	UniProt ID
	Human	<u>4233;</u>	<u>P08581;</u>
	Mouse		<u>P16056;</u>
	Rat	<u>24553;</u>	<u>P97523;</u>
Cellular Localization	Membrane; Single-pass type I membrane protein.; [Isoform 3]: Secreted.		
Tissue specificity	Expressed in normal hepatocytes as well as in epithelial cells lining the stomach, the small and the large intestine. Found also in basal keratinocytes of esophagus and skin. High levels are found in liver, gastrointestinal tract, thyroid and kidney. Also present in the brain. Expressed in metaphyseal bone (at protein level) (PubMed:26637977).		
Function	Expressed in metaphyseal bone (at protein level) (PubMed:26637977). Catalytic activity:ATP + a [protein]-L-tyrosine = ADP + a [protein]-L-tyrosine phosphate.,Disease:Activation of MET after rearrangement with the TPR gene produces an oncogenic protein.,Disease:Defects in MET are a cause of hepatocellular carcinoma (HCC) [MIM:114550].,Disease:Defects in MET are a cause of hereditary papillary renal carcinoma (HPRC) [MIM:605074]; also known as papillary renal cell carcinoma 2 (RCCP2). HPRC is a form of inherited kidney cancer characterized by a predisposition to develop multiple, bilateral papillary renal tumors. The pattern of inheritance is consistent with autosomal dominant transmission with reduced penetrance.,Disease:Defects in MET may be associated with gastric cancer.,Disease:Genetic variations in MET may be associated with susceptibility to autism type 9 (AUTS9) [MIM:611015]. Autism is a neurodevelopmental disorder characterized by disturbance in language, perception and socialization. The disorder is classically defined by a triad of limited or absent verbal communication, a lack of reciprocal social interaction or responsiveness, and restricted, stereotypical, and ritualized patterns of interests and behavior.,Domain:The kinase domain is involved in SPSB1 binding,.Function:Receptor for hepatocyte growth factor and scatter factor. Has a tyrosine- protein kinase activity. Functions in cell proliferation, scattering, morphogenesis and survival.,online information:C-MET entry,similarity:Contains 1 protein kinase domain.,similarity:Contains 1 Sema domain.,similarity:Contains 3 IPT/TIG domains.,subunit:Heterodimer formed of an alpha chain (50 kDa) and a beta chain (145 kDa) which are disulfide linked. Binds PLXNB1 and GRB2. Interacts with SPSB1, SPSB2 and SPSB4 (By similarity). Interacts with INPP5D/SHIP1. When phosphorylated at Tyr-1356, interacts with INPPL1/SHIP2. Interacts with RANBP9 and RANBP10, as well as SPSB1, SPSB2, SPSB3 and SPSB4. SPSB1 binding occurs in the presence and in the absence of HGF, how		

Validation Data





Observed band size: 90kDa

Immunofluorescence analysis of HEK293. Picture A: Met (Phospho Tyr1234/1235) antibody (red). Picture B: DAPI (blue). Picture C: Merge of A+B

Contact information

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Please scan the QR code to access additional product information: Met (Phospho Tyr1234/1235) (PT0745R) PT® Rabbit mAb

For Research Use Only. Not for Use in Diagnostic Procedures.

Antibody | ELISA Kits | Protein | Reagents