

Catenin-β (Phospho Thr41/Ser45) Rabbit pAb

CatalogNo: YP0048

Key Features

Host Species			
 Rabbit 			

Reactivity

Human,Mouse,Rat

ApplicationsWB,ELISA

MW • 92kD (Observed)

Isotype • lgG

Recommended Dilution Ratios

WB 1:500-1:2000 ELISA 1:20000 Not yet tested in other applications.

Storage

Storage*-15°C to -25°C/1 year(Do not lower than -25°C)FormulationLiquid in PBS containing 50% glycerol, 0.5% BSA and 0.02% sodium azide.

Basic Information

Clonality Polyclonal

Immunogen Information

Immunogen The antiserum was produced against synthesized peptide derived from human Cateninbeta around the phosphorylation site of Thr41/Ser45. AA range:11-60

Specificity This antibody detects endogenous levels of beta-catenin only when phosphorylated at Thr41 or Ser45. This antibody does not recognize β-catenin phosphorylated at other sites. The name of modified sites may be influenced by many factors, such as species (the modified site was not originally found in human samples) and the change of protein sequence (the previous protein sequence is incomplete, and the protein sequence may be prolonged with the development of protein sequencing technology). When naming, we will use the "numbers" in historical reference to keep the sites consistent with the reports. The antibody binds to the following modification sequence (lowercase letters are modification sites):ATtTAPsLS

Target Information

CTNNB1 CTNNB OK/SW-cl.35 PRO2286

Protein Name

Cellular Localizatio

Tissue spe

Catenin-β:b-catenin;Beta catenin;Beta-catenin;Cadherin associated protein;Catenin (cadherin associated protein), beta 1, 88 kDa;Catenin beta 1;Catenin beta-1;CATNB;CHBCAT;CTNB1_HUMAN;CTNNB;CTNNB1;DKFZp686D02253;FLJ25606;FLJ37923;OTTHUMP00000162082;OTTHUMP0000165222;OTTHUMP00000165223;OTTHUMP00000209288;OTTHUMP00000209289

	Organism	Gene ID	UniProt ID	
	Human	<u>1499;</u>	<u>P35222;</u>	
	Mouse	12387;	<u>Q02248;</u>	
	Rat	<u>84353;</u>	<u>09WU82;</u>	
	Cytoplasm. Nucleus . Cytoplasm, cytoskeleton . Cell junction, adherens junction . Cell junction . Cell membrane . Cytoplasm, cytoskeleton, microtubule organizing center, centrosome. Cytoplasm, cytoskeleton, spindle pole. Cell junction, synapse . Cytoplasm, cytoskeleton, cilium basal body . Colocalized with RAPGEF2 and TJP1 at cell-cell contacts (By similarity). Cytoplasmic when it is unstabilized (high level of phosphorylation) or bound to CDH1. Translocates to the nucleus when it is stabilized (low level of phosphorylation). Interaction with GLIS2 and MUC1 promotes nuclear translocation. Interaction with EMD inhibits nuclear localization. The majority of beta-catenin is localized to the cell membrane. In interphase, colocalizes with CROCC between CEP250 punct at the proximal end of centrioles, and this localization is dependent on CROCC and CEP250. In mitosis, when NEX2 activity increases, it localizes to centrosomes at spindle poles independent of CROCC. Colocalizes with CCMS in the cell-cell contacts and plasma membrane of undifferentiated and differentiated neuroblastoma cells. Interaction with FAM53B promotes translocation to the nucleus (PubMed:25183871)			
pecificity	Expressed in several hair follicle cell types: basal and peripheral ma tissues (at protein level) (PubMed:29367600).	trix cells, and cells of the outer and inner root sheaths. Expressed	d in colon. Present in cortical neurons (at protein level). Expressed in breast cancer	

tissues (at protein level) (PubMed:29367600). Disease:A chromosomal rearrangement involving CTNNB1 may be a cause of salivary gland pleiomorphic adenomas (PA) [181030]. Pleiomorphic adenomas are the most common benign epithelial tumors of the salivary gland. Translocation t13;8)(p21;q12) with PLAG1., Disease:Activating mutations in CTNNB1 have oncogenic activity resulting in tumor development. Somatic mutations are found in various tumor types, including colon cancers, ovarian and prostate carcinomas, hepatoblastoma (HB), hepatocellular carcinoma (HCC). HBs are malignant embryonal tumors mainly affecting young children in the first three years of life.,Disease:Defects in CTNNB1 are a cause of medulloblastoma (MDB) (MIN:15525). MDB is a malignant, invasive embryonal tumor of the cerebellum with a preferential manifestation in children.,Disease:Defects in CTNNB1 are a cause of joinmatrixoma (PTR) (MIN:132600); a common benign skin tumor,.Disease:Defects in CTNNB1 are a sociated with colrectal cancer (CRC) (MIN:145000). Disease:Defects in CTNNB1 are a sociated with ovarian cancer (RCN) (MIN:145000). Joint cancer is the leading cause of death from gynecologic malignancy. It is characterized by advanced presentation with loco-regional dissemination in the peritoneal cavity and the rare incidence of visceral metastases. These typical features relate to the disease, which is a principal determinant of outcome, Function:Thoveld in the regulation of cell adhesion and in signal transduction through the Wnt pathway, online information:Beta-catenin entry, PTM:EGF stimulates tyrosine phosphorylation on Tyr-654 decreases CDH1 binding and enhances TBP binding,.PTM:Hosphorylation by GSXB requires prior phosphorylation of Ser-45 by another kinase. Phosphorylation proceeds then from Thr-41 to Ser-33, PTM:Ubiquitinated by a E3 ubiquitination (HB1LX) (Probabile). Its ubiquitination berciton with dis2 and MUC1 promotes nuclear translocation.subunit:Two separate pools are found in the cytoplasm: one is PSENL/Cadherin/Cadherin/Cadher Function (via the cytoplasmic juxtamembrane domain)..tissue specificity:Expressed in several hair follicle cell types: basal and peripheral matrix cells, and cells of the outer and inner root sheats. Expressed in colon.

Validation Data

Contact information

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Please scan the QR code to access additional product information: Catenin-β (Phospho Thr41/Ser45) Rabbit pAb

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

Antibody | ELISA Kits | Protein | Reagents