

Claudin-4 (phospho Tyr208) Polyclonal Antibody

Catalog No: YP1088

Reactivity: Human; Rat; Mouse;

Applications: IHC;IF;ELISA

Target: Claudin-4

Fields: >>Cell adhesion molecules;>>Tight junction;>>Leukocyte transendothelial

migration;>>Pathogenic Escherichia coli infection;>>Hepatitis C

Gene Name: CLDN4

Protein Name: Claudin-4

Human Gene Id: 1364

Human Swiss Prot

No:

Mouse Swiss Prot

No:

Immunogen: The antiserum was produced against synthesized peptide derived from human

Claudin 4 around the phosphorylation site of Tyr208. AA range:160-209

Specificity: Phospho-Claudin-4 (Y208) Polyclonal Antibody detects endogenous levels of

Claudin-4 protein only when phosphorylated at Y208.

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

Source: Polyclonal, Rabbit, IgG

O14493

O35054

Dilution: IHC 1:100 - 1:300. ELISA: 1:5000.. IF 1:50-200

Purification: The antibody was affinity-purified from rabbit antiserum by affinity-

chromatography using epitope-specific immunogen.

Concentration: 1 mg/ml

1/2



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

Molecularweight: 22kD

Cell Pathway: Cell adhesion molecules (CAMs); Tight junction; Leukocyte transendothelial

migration;

Background: The protein encoded by this intronless gene belongs to the claudin family.

Claudins are integral membrane proteins that are components of the epithelial cell

tight junctions, which regulate movement of solutes and ions through the paracellular space. This protein is a high-affinity receptor for Clostridium

perfringens enterotoxin (CPE) and may play a role in internal organ development and function during pre- and postnatal life. This gene is deleted in Williams-Beuren syndrome, a neurodevelopmental disorder affecting multiple systems.

[provided by RefSeq, Sep 2013],

Function: disease: Haploin sufficiency of CLDN4 may be the cause of certain

cardiovascular and musculo-skeletal abnormalities observed in Williams-Beuren syndrome (WBS), a rare developmental disorder. It is a contiguous gene deletion syndrome involving genes from chromosome band 7g11.23..function:Plays a

major role in tight junction-specific obliteration of the intercellular

space., similarity: Belongs to the claudin family., subunit: Directly interacts with

TJP1/ZO-1, TJP2/ZO-2 and TJP3/ZO-3.,

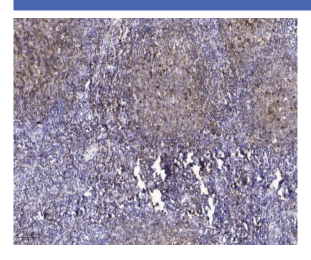
Subcellular Location:

Cell junction, tight junction. Cell membrane; Multi-pass membrane protein.

CLDN4 is required for tight junction localization in the kidney. .

Expression : Colon, Fetal brain, Trachea,

Products Images



Immunohistochemical analysis of paraffin-embedded human tonsil. 1, Antibody was diluted at 1:200(4° overnight). 2, Tris-EDTA,pH9.0 was used for antigen retrieval. 3,Secondary antibody was diluted at 1:200(room temperature, 45min).