

Collagen IV Rabbit pAb

Collagen IV Rabbit pAb Catalog # AP94831

Product Information

Application WB, IHC-P, IHC-F, IF, E

Primary Accession
Reactivity
Rat, Human
Host
Clonality
Polyclonal
Calculated MW
Physical State
P02463
Rat, Human
Rabbit
Polyclonal
160679
Liquid

Immunogen KLH conjugated synthetic peptide derived from human Collagen alpha-1(IV)

chain

Epitope Specificity 1571-1669/1669

Isotype IgG

Purity affinity purified by Protein A

Buffer SUBCELLULAR LOCATION

SIMILARITY

0.01M TBS (pH7.4) with 1% BSA, 0.02% Proclin300 and 50% Glycerol. Secreted, extracellular space, extracellular matrix, basement membrane. Contains 1 FAD-binding FR-type domain.Contains 1 ferric oxidoreductase

domain.

SUBUNIT There are six type IV collagen isoforms, alpha 1(IV)-alpha 6(IV), each of which

in the development of the cerebral ventricles.

can form a triple helix structure with 2 other chains to generate type IV

collagen network.

Post-translational modifications

Lysines at the third position of the tripeptide repeating unit (G-X-Y) are hydroxylated in all cases and bind carbohydrates. Prolines at the third position of the tripeptide repeating unit (G-X-Y) are hydroxylated in some or all of the chains. Type IV collagens contain numerous cysteine residues which are involved in inter- and intramolecular disulfide bonding. 12 of these, located in the NC1 domain, are conserved in all known type IV collagens. The trimeric structure of the NC1 domains is stabilized by covalent bonds between Lys and Met residues. Proteolytic processing produces the C-terminal NC1 peptide, arresten.

DISEASE

Defects in COL4A1 are a cause of brain small vessel disease with hemorrhage (BSVDH) [MIM:607595]. Brain small vessel diseases underlie 20 to 30 percent of ischemic strokes and a larger proportion of intracerebral hemorrhages. Inheritance is autosomal dominant. Defects in COL4A1 are the cause of hereditary angiopathy with nephropathy aneurysms and muscle cramps (HANAC) [MIM:611773]. The clinical renal manifestations include hematuria and bilateral large cysts. Histologic analysis revealed complex basement membrane defects in kidney and skin. The systemic angiopathy appears to affect both small vessels and large arteries. Defects in COL4A1 are a cause of familial porencephaly (POREN1) [MIM:175780]. Porencephaly is a term used for any cavitation or cerebrospinal fluid-filled cyst in the brain. Porencephaly type 1 is usually unilateral and results from focal destructive lesions such as fetal vascular occlusion or birth trauma. Type 2, or schizencephalic porencephaly, is usually symmetric and represents a primary defect or arrest

Important Note This product as supplied is intended for research use only, not for use in

human, therapeutic or diagnostic applications.

Background Descriptions This gene encodes the major type IV alpha collagen chain of basement

membranes. Like the other members of the type IV collagen gene family, this gene is organized in a head-to-head conformation with another type IV collagen gene so that each gene pair shares a common promoter. [provided

by RefSeq, Jul 2008]

Additional Information

Gene ID 12826

Other Names Collagen alpha-1(IV) chain, Arresten, Col4a1 {ECO:0000312|MGI:MGI:88454}

Target/Specificity Highly expressed in placenta.

Dilution WB=1:500-2000,IHC-P=1:100-500,IHC-F=1:100-500,IF=1:100-500,Flow-Cyt=2ug

/Test,ELISA=1:5000-10000

Format 0.01M TBS(pH7.4) with 1% BSA, 0.09% (W/V) sodium azide and 50% Glyce

Storage Store at -20 °C for one year. Avoid repeated freeze/thaw cycles. When

reconstituted in sterile pH 7.4 0.01M PBS or diluent of antibody the antibody

is stable for at least two weeks at 2-4 °C.

Protein Information

Name Col4a1 {ECO:0000312 | MGI:MGI:88454}

Function Type IV collagen is the major structural component of glomerular basement

membranes (GBM), forming a 'chicken-wire' meshwork together with

laminins, proteoglycans and entactin/nidogen.

Cellular Location Secreted, extracellular space, extracellular matrix, basement membrane

Tissue Location Detected in the basement membrane of the cornea (at protein level).

Background

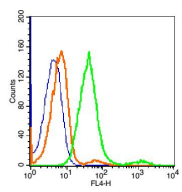
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Images

Blank control: Hepg2 Cells(blue). Primary Antibody: Rabbit Anti-Collagen IV/AF647 Conjugated antibody (AP94831), Dilution: 1 μ g in 100 μ L 1X PBS containing

0.5% BSA; Isotype Control Antibody: Rabbit

IgG/AF647(orange), used under the same conditions.



Please note: All products are 'FOR RESEARCH USE ONLY. NOT FOR USE IN DIAGNOSTIC OR THERAPEUTIC PROCEDURES'.