

## Collagen IV α1 (Cleaved-Gly173) rabbit pAb

Cat No.: ES19975

For research use only

## Overview

Product Name Collagen IV α1 (Cleaved-Gly173) rabbit pAb

Host species Rabbit
Applications WB; ELISA
Species Cross-Reactivity Human; Mouse

Recommended dilutions WB 1:1000-2000 ELISA 1:5000-20000

Immunogen Synthesized peptide derived from human Collagen

IV α1 (Cleaved-Gly173)

**Specificity** This antibody detects endogenous levels of

Human, Mouse Collagen IV  $\alpha 1$  (Cleaved-Gly173, protein was cleaved amino acid sequence between

172-173)

Formulation Liquid in PBS containing 50% glycerol, 0.5% BSA and

0.02% sodium azide.

**Storage** Store at  $-20^{\circ}$ C. Avoid repeated freeze-thaw cycles.

Protein Name Collagen IV α1 (Cleaved-Gly173)

Gene Name COL4A1

**Cellular localization** Secreted, extracellular space, extracellular matrix,

basement membrane.

**Purification** The antibody was affinity-purified from rabbit

antiserum by affinity-chromatography using

epitope-specific immunogen.

ClonalityPolyclonalConcentration1 mg/mlObserved band18 185kDHuman Gene ID1282Human Swiss-Prot NumberP02462

Background

Alternative Names Collagen alpha-1(IV) chain [Cleaved into: Arresten]

disease:Defects in COL4A1 are a cause of brain small vessel disease with hemorrhage [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., disease: Defects in COL4A1 are a cause of



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porencephaly type 1 [MIM:175780]; also known as encephaloclastic porencephaly. 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. Inheritance is autosomal dominant., disease: 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., domain: Alpha chains of type IV collagen have a non-collagenous domain (NC1) at their C-terminus, frequent interruptions of the G-X-Y repeats in the long central triple-helical domain (which may cause flexibility in the triple helix), and a short N-terminal triple-helical 7S domain., 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. Potently inhibits endothelial cell proliferation and angiogenesis. Inhibits angiogenesis potentially via mechanisms involving cell surface proteoglycans and the alpha and beta integrins of endothelial cells.,PTM:Lysines at the third position of the tripeptide repeating unit (G-X-Y) are hydroxylated in all cases and bind carbohydrates., PTM: Prolines at the third position of the tripeptide repeating unit (G-X-Y) are hydroxylated in some or all of the chains.,PTM:The trimeric structure of the NC1 domains may be stabilized by covalent bonds between Lys and Met residues., PTM: 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., similarity: Belongs to the



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type IV collagen family.,similarity:Contains 1 collagen IV NC1 (C-terminal non-collagenous) domain.,subunit:There are six type IV collagen isoforms, alpha 1(IV)-alpha 6(IV), each of which can form a triple helix structure with 2 other chains to generate type IV collagen network.,tissue specificity:Highly expressed in placenta.,



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