



Collagen XI α1 Monoclonal Antibody

Catalog No	BYmab-10810
lsotype	lgG
Reactivity	Human;Mouse;Rat
Applications	WB
Gene Name	COL11A1 COLL6
Protein Name	Collagen alpha-1(XI) chain
Immunogen	Synthesized peptide derived from human Collagen XI α 1 Monoclonal
Specificity	This antibody detects endogenous levels of Collagen XI α 1.
Formulation	Liquid in PBS containing 50% glycerol, 0.5% BSA and 0.02% sodium azide.
Source	Monoclonal, Mouse,IgG
Purification	The antibody was affinity-purified from mouse antiserum by affinity-chromatography using epitope-specific immunogen.
Dilution	WB 1:500-2000
Concentration	1 mg/ml
Purity	≥90%
Storage Stability	-20°C/1 year
Synonyms	Collagen alpha-1(XI) chain
Observed Band	180kD
Cell Pathway	Secreted, extracellular space, extracellular matrix .
Tissue Specificity	Cartilage, placenta and some tumor or virally transformed cell lines. Isoforms using exon IIA or IIB are found in the cartilage while isoforms using only exon IIB are found in the tendon.
Function	alternative products:Additional isoforms seem to exist. There is alternative usage of exon IIA or exon IIB. Transcripts containing exon IIA or IIB are present in cartilage, but exon IIB is preferentially utilized in transcripts from tendon,disease:Defects in COL11A1 are the cause of Marshall syndrome [MIM:154780]. It is an autosomal dominant disorder with ocular, orofacial, auditory and skeletal manifestations. It shares several features with Stickler syndrome, such as midfacial hypoplasia, high myopia, and sensorineural-hearing deficit.,disease:Defects in COL11A1 are the cause of Stickler syndrome type 2 (STL2) [MIM:604841]; also known as Stickler syndrome vitreous type 2. STL2 is an autosomal dominant form of Stickler syndrome, an inherited disorder that associates ocular signs with more or less complete forms of Pierre Robin
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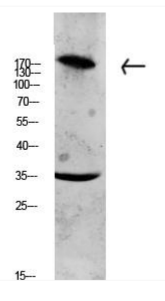
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collagen is a heterotrime modified alpha 1 type II Stickler syndrome and w in this gene is also asso	chain(COL11A1) Homo sapiens This gene encodes ains of type XI collagen, a minor fibrillar collagen. Type XI er but the third alpha chain is a post-translationally chain. Mutations in this gene are associated with type II vith Marshall syndrome. A single-nucleotide polymorphism ciated with susceptibility to lumbar disc herniation. the have been identified for this gene. [provided by
matters needingAvoid repeated freezingattention	and thawing!
Usage suggestions This product can be use more information, please	d in immunological reaction related experiments. For e consult technical personnel.

Products Images



Western Blot analysis of various cells using Collagen XI α 1 Monoclonal Antibody

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