



COL1A2 Monoclonal Antibody

| Catalog No | BYmab-16977 |
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| lsotype | lgG |
| Reactivity | Human;Mouse;Rat |
| Applications | WB |
| Gene Name | COL1A2 |
| Protein Name | Collagen alpha-2(I) chain |
| Immunogen | The antiserum was produced against synthesized peptide derived from human Collagen I. AA range:1-50 |
| Specificity | COL1A2 Monoclonal Antibody detects endogenous levels of COL1A2 protein. |
| 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 | COL1A2; Collagen alpha-2(I) chain; Alpha-2 type I collagen |
| Observed Band | 200kD |
| Cell Pathway | Secreted, extracellular space, extracellular matrix . |
| Tissue Specificity | Forms the fibrils of tendon, ligaments and bones. In bones the fibrils are mineralized with calcium hydroxyapatite. |
| Function | disease:A chromosomal rearrangement involving COL1A2 may be a cause of lipoblastomas, which are benign tumors resulting from transformation of adipocytes, usually diagnosed in children. Translocation t(7;8)(p22;q13) with PLAG1.,disease:Defects in COL1A2 are a cause of osteogenesis imperfecta type I (OI-I) [MIM:166200]. OI-I is a dominantly inherited serious newborn disease characterized by bone fragility, normal stature, little or no deformity, blue sclerae and hearing loss in 50% of families. Dentinogenesis imperfecta is rare and may distinguish a subset of OI type I (formation of dentine).,disease:Defects in COL1A2 are a cause of osteogenesis imperfects in COL1A2 are a cause of osteogenesis imperfecta type II (OI-II) [MIM:166210]; also |
| | serious newborn disease that diffusely affects bone. Infants are born with multiple |

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网址: www.njbybio.com 官方热线: 025-5229-8998 监督电话: 15950492658





fractures, which lead to shortening

| Background | This gene encodes the pro-alpha2 chain of type I collagen whose triple helix comprises two alpha1 chains and one alpha2 chain. Type I is a fibril-forming collagen found in most connective tissues and is abundant in bone, cornea, dermis and tendon. Mutations in this gene are associated with osteogenesis imperfecta types I-IV, Ehlers-Danlos syndrome type VIIB, recessive Ehlers-Danlos syndrome Classical type, idiopathic osteoporosis, and atypical Marfan syndrome. Symptoms associated with mutations in this gene, however, tend to be less severe than mutations in the gene for the alpha1 chain of type I collagen (COL1A1) reflecting the different role of alpha2 chains in matrix integrity. Three transcripts, resulting from the use of alternate polyadenylation signals, have been identified for this gene. [provided by R. Dalgleish, Feb 2008], |
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| matters needing attention | Avoid repeated freezing and thawing! |
| Usage suggestions | This product can be used in immunological reaction related experiments. For more information, please consult technical personnel. |

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