

Human Recombinant Carbonic Anhydrase 2

CATALOG #:	6390-100	100 µg
ALTERNATE NAMES:	CA-II, CAC, Carbonic anhydrase 2, Carbonate dehydratase 2, can, cynT2, yadF, b0126, JW0122, Carbonic Anhydrase II.	
SOURCE:	E.Coli	
PURITY:	> 95% by SDS - PAGE	
MOL. WEIGHT:	29.2 kDa (260 aa, 1-260 aa)	
FORMULATION:	1 mg/ml solution in 20 mM Tris-HCl buffer (pH 8.0) containing 50 mM NaCl, 1 mM DTT and 10% glycerol.	

STORAGE CONDITIONS:

Can be stored at 4 °C short term (1-2 weeks). For long term storage, aliquot and store at -20 °C or -70 °C. Avoid repeated freezing and thawing cycles.

DESCRIPTION:

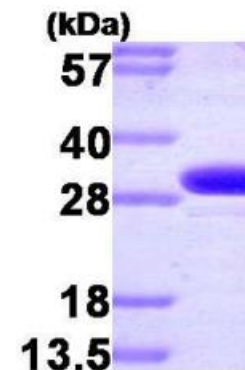
The enzyme Carbonic anhydrase 2 (CA2) is a part of the enzyme family that catalyses rapid inter-conversion of carbon dioxide & water to bicarbonate, carbonic acid and protons ($\text{CO}_2 + \text{H}_2\text{O} \rightarrow \text{HCO}_3 + \text{H}$), a reaction that occurs rather slowly in the absence of a catalyst. The majority of carbonic anhydrases enclose a zinc ion in their active site and therefore is classified as metalloenzymes. The most important function of Carbonic anhydrase is known to preserve acid-base balance in blood and other tissues, and to help transport carbon dioxide of tissues. Carbonic anhydrases have been found in all kingdoms of life. Mammalian carbonic anhydrase is monomeric and belongs to the alpha class. Mutations in the CA2 gene result in the CA2 deficiency syndrome, an autosomal recessive disorder that produces osteoporosis, renal tubular acidosis and cerebral calcification.

AMINO ACID SEQUENCE:

MSHHWGYGKH NGPEHWHKDF PIAKGERQSP VDIDHTAKY DPSLKPLSVS
YDQATSLRIL NNGHAFNVEF DDSQDKAVLK GGPLDGTYRL IQFHFHWGSL
DGQGSEHTVD KKKYAAELHL VHWNTKYGDF GKAVQQPDGL AVLGIFLKV
SAKPLGQKVV DVLDSIKTKG KSADFTNFDP RGLLPESLDY WTPGSLTTP PLLECVTWIV
LKEPISVSSE QVLKFRKLNFGEGEPEELM VDNWRPAQPL KNRQIKASFK

BIOLOGICAL ACTIVITY:

Specific activity is > 7000 pmoles/min/µg and is defined as the amount of enzyme that hydrolyzes 1.0 pmole of 4-nitrophenyl acetate to 4-nitrophenol per minute at pH 8.0 at 37 °C.



15% SDS-PAGE (3ug)

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RELATED PRODUCTS:

- MMP-1, human recombinant (Cat. No. 7781-10, 50, 1000)
- MMP-2, human recombinant (Cat. No. 7782-10, 50, 1000)
- MMP-3, human recombinant (Cat. No. 7783-10, 50, 1000)
- MMP-9, human recombinant (Cat. No. 7789-10, 50, 1000)

FOR RESEARCH USE ONLY! Not to be used in humans.