

## Hexokinase 1 mouse mAb

YM1282 Catalog No:

Human:Mouse:Rat Reactivity:

**Applications: WB** 

HXK I Target:

Fields: >>Glycolysis / Gluconeogenesis;>>Fructose and mannose

metabolism;>>Galactose metabolism;>>Starch and sucrose

metabolism;>>Amino sugar and nucleotide sugar metabolism;>>Neomycin, kanamycin and gentamicin biosynthesis;>>Metabolic pathways;>>Carbon

metabolism;>>Biosynthesis of nucleotide sugars;>>HIF-1 signaling

pathway;>>Insulin signaling pathway;>>Type II diabetes mellitus;>>Carbohydrate digestion and absorption;>>Shigellosis;>>Central carbon metabolism in cancer

Gene Name: hk1

**Human Gene Id:** 3098

**Human Swiss Prot** 

No:

**Mouse Swiss Prot** 

No:

P17710

Purified recombinant human Hexokinase 1 protein fragments expressed in E.coli Immunogen:

**Specificity:** This antibody detects endogenous levels of Hexokinase 1 and does not cross-

react with Hexokinase 2 and other proteins.

Formulation: Liquid in PBS containing 50% glycerol, 0.5% BSA and 0.02% sodium azide.

Source: Monoclonal, Mouse

**Dilution:** wb 1:1000

**Purification:** The antibody was affinity-purified from mouse ascites by affinity-

chromatography using epitope-specific immunogen.

1 mg/ml

P19367



**Storaget Stability:** -15°C to -25°C/1 year(Do not lower than -25°C)

Observed Band: 102kD

**Cell Pathway:** Glycolysis / Gluconeogenesis; Fructose and mannose metabolism; Galactose

metabolism;Starch and sucrose metabolism;Amino sugar and nucleotide sugar

metabolism;Insulin\_Receptor;Type II diabetes mellitus;

**Background :** Hexokinases phosphorylate glucose to produce glucose-6-phosphate, the first

step in most glucose metabolism pathways. This gene encodes a ubiquitous form of hexokinase which localizes to the outer membrane of mitochondria. Mutations in this gene have been associated with hemolytic anemia due to hexokinase deficiency. Alternative splicing of this gene results in several transcript variants which encode different isoforms, some of which are tissue-specific. [provided by

RefSeq, Apr 2016],

**Function:** catalytic activity:ATP + D-hexose = ADP + D-hexose

6-phosphate., disease: Defects in HK1 are the cause of hexokinase deficiency [MIM:235700]. Hexokinase deficiency is a rare autosomal recessive disease with nonspherocytic hemolytic anemia as the predominant clinical feature., domain: The N- and C-terminal halves of this hexokinase show extensive sequence similarity to

each other. The catalytic activity is associated with the C-terminus while

regulatory function is associated with the N-terminus., enzyme

regulation:Hexokinase is an allosteric enzyme inhibited by its product Glc-6-P.,miscellaneous:In vertebrates there are four major glucose-phosphorylating isoenzymes, designated hexokinase I, II, III and IV (glucokinase).,online information:Hexokinase entry,pathway:Carbohydrate

metabolism; hexose metabolism., similarity: Belongs to the hexokinase

family., subcellular location: Its hydrophobic N-ter

Subcellular Location:

Mitochondrion outer membrane; Peripheral membrane protein. Cytoplasm, cytosol. The mitochondrial-binding peptide (MBP) region promotes association

with the mitochondrial outer membrane (Probable). Dissociates from the mitochondrial outer membrane following inhibition by N-acetyl-D-glucosamine,

leading to relocation to the cytosol (PubMed:27374331). .

**Expression :** Isoform 2: Erythrocyte specific (Ref.6). Isoform 3: Testis-specific

(PubMed:10978502). Isoform 4: Testis-specific (PubMed:10978502).

**Sort :** 7332

No4: 1

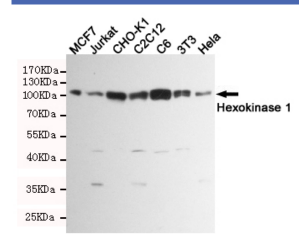
Host: Mouse

Modifications: Unmodified

2/3



## **Products Images**



Western blot detection of Hexokinase 1 in MCF7, Jurkat, CHO-K1, C2C12, C6, 3T3 and Hela cell lysates using Hexokinase 1 mouse mAb (1:1000 diluted). Predicted band size:102KDa. Observed band size:102KDa.