abcam

Product datasheet

Recombinant Human PMM2 protein ab99391

1 Image

Description

Product name Recombinant Human PMM2 protein

Purity > 90 % SDS-PAGE.

ab99391 is purified using conventional chromatography techniques.

Expression system Escherichia coli

Accession O15305

Protein length Full length protein

Animal free No

Nature Recombinant

Species Human

Sequence MGSSHHHHHHSSGLVPRGSHMAAPGPALCLFDVDGTL

TAPRQKITKEMDD

FLQKLRQKIKIGVVGGSDFEKVQEQLGNDVVEKYDYVFPE

NGLVAYKDGK

LLCRQNIQSHLGEALIQDLINYCLSYIAKIKLPKKRGTFIEFRN

GMLNVS

PIGRSCSQEERIEFYELDKKENIRQKFVADLRKEFAGKGLT

FSIGGQISF

DVFPDGWDKRYCLRHVENDGYKTIYFFGDKTMPGGNDHE

IFTDPRTMGYS VTAPEDTRRICELLFS

Predicted molecular weight 30 kDa including tags

Amino acids 1 to 246

Tags His tag N-Terminus

Specifications

Our **Abpromise guarantee** covers the use of **ab99391** in the following tested applications.

The application notes include recommended starting dilutions; optimal dilutions/concentrations should be determined by the end user.

Applications SDS-PAGE

Mass Spectrometry

Mass spectrometry MALDI-TOF

Form Liquid

1

Preparation and Storage

Stability and Storage

Shipped at 4°C. Upon delivery aliquot and store at -20°C or -80°C. Avoid repeated freeze / thaw cycles.

pH: 8.00

Constituents: 0.0154% DTT, 0.316% Tris HCl, 10% Glycerol (glycerin, glycerine), 0.58% Sodium chloride

General Info

Function Involved in the synthesis of the GDP-mannose and dolichol-phosphate-mannose required for a

number of critical mannosyl transfer reactions.

Pathway Nucleotide-sugar biosynthesis; GDP-alpha-D-mannose biosynthesis; alpha-D-mannose 1-

phosphate from D-fructose 6-phosphate: step 2/2.

Involvement in diseaseDefects in PMM2 are the cause of congenital disorder of glycosylation type 1A (CDG1A)

[MIM:212065]; also known as carbohydrate-deficient glycoprotein syndrome type la (CDGS1A) or

Jaeken syndrome. Congenital disorders of glycosylation are metabolic deficiencies in

glycoprotein biosynthesis that usually cause severe mental and psychomotor retardation. They are characterized by under-glycosylated serum glycoproteins. CDG1A is an autosomal recessive

disorder characterized by a severe encephalopathy with axial hypotonia, abnormal eye

movement, and pronounced psychomotor retardation, as well as peripheral neuropathy, cerebellar

hypoplasia, and retinitis pigmentosa. Patients show a peculiar distribution of subcutaneous fat,

nipple retraction, and hypogonadism.

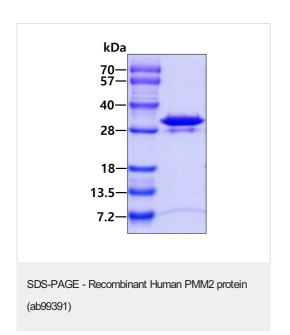
Sequence similarities

Belongs to the eukaryotic PMM family.

Cellular localization

Cytoplasm.

Images



15% SDS-PAGE analysis of 3µg ab99391.

Please note: All products are "FOR RESEARCH USE ONLY. NOT FOR USE IN DIAGNOSTIC PROCEDURES"

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