abcam

Product datasheet

Anti-ALG2 antibody [EPR12006(2)(B)] ab178697



RabMAb

2 Images

Overview

Product name Anti-ALG2 antibody [EPR12006(2)(B)]

Description Rabbit monoclonal [EPR12006(2)(B)] to ALG2

Host species Rabbit

Tested applications Suitable for: WB

Unsuitable for: Flow Cyt,ICC/IF,IHC-P or IP

Species reactivity Reacts with: Human

Immunogen Synthetic peptide within Human ALG2 aa 1-100 (Cysteine residue). The exact sequence is

proprietary.

Database link: Q9H553

Positive control HepG2, HeLa, A431 and Human fetal kidney lysates.

General notes This product is a recombinant monoclonal antibody, which offers several advantages including:

- High batch-to-batch consistency and reproducibility

Improved sensitivity and specificityLong-term security of supply

- Animal-free production

For more information see here.

Our RabMAb[®] technology is a patented hybridoma-based technology for making rabbit monoclonal antibodies. For details on our patents, please refer to **RabMAb**[®] **patents**.

Mouse, Rat: We have preliminary internal testing data to indicate this antibody may not react with

these species. Please contact us for more information.

Properties

Form Liquid

Storage instructions Shipped at 4°C. Store at +4°C short term (1-2 weeks). Upon delivery aliquot. Store at -20°C long

term. Avoid freeze / thaw cycle.

Storage buffer pH: 7.20

Preservative: 0.01% Sodium azide

Constituents: 9% PBS, 40% Glycerol (glycerin, glycerine), 0.05% BSA, 50% Tissue culture

supernatant

1

Purity Tissue culture supernatant

Clonality Monoclonal

Clone number EPR12006(2)(B)

Isotype IgG

Applications

The Abpromise guarantee Our Abpromise guarantee covers the use of ab178697 in the following tested applications.

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

Application	Abreviews	Notes
WB		1/10000 - 1/50000. Predicted molecular weight: 47 kDa.

Application notes Is unsuitable for Flow Cyt,ICC/IF,IHC-P or IP.

Target

Function Mannosylates Man(2)GlcNAc(2)-dolichol diphosphate and Man(1)GlcNAc(2)-dolichol diphosphate

to form Man(3)GlcNAc(2)-dolichol diphosphate.

Pathway Protein modification; protein glycosylation.

Involvement in diseaseDefects in ALG2 are the cause of congenital disorder of glycosylation type 1I (CDG1I)

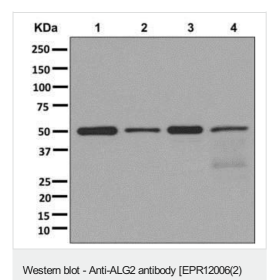
[MIM:607906]. CDGs are a family of severe inherited diseases caused by a defect in protein N-glycosylation. They are characterized by under-glycosylated serum proteins. These multisystem disorders present with a wide variety of clinical features, such as disorders of the nervous system development, psychomotor retardation, dysmorphic features, hypotonia, coagulation disorders, and immunodeficiency. The broad spectrum of features reflects the critical role of N-glycoproteins

during embryonic development, differentiation, and maintenance of cell functions.

Sequence similarities Belongs to the glycosyltransferase 1 family.

Cellular localization Membrane.

Images



(B)] (ab178697)

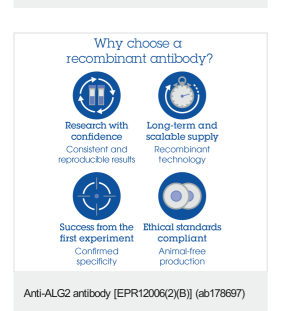
All lanes : Anti-ALG2 antibody [EPR12006(2)(B)] (ab178697) at 1/10000 dilution

Lane 1 : HepG2 lysate Lane 2 : HeLa lysate Lane 3 : A431 lysate

Lane 4: Human fetal kidney lysate

Lysates/proteins at 10 µg per lane.

Predicted band size: 47 kDa



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

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