

Dymeclin Polyclonal Antibody

Purified Rabbit Polyclonal Antibody (Pab) Catalog # AP55044

Product Information

Application WB, IHC-P, IHC-F, IF, ICC, E

Primary Accession <u>Q7RTS9</u>

Reactivity Rat, Dog, Bovine

Host Rabbit
Clonality Polyclonal
Calculated MW 75935
Physical State Liquid

Immunogen KLH conjugated synthetic peptide derived from human Dymeclin

Epitope Specificity 151-250/669

Isotype IgG

Purity affinity purified by Protein A

Buffer 0.01M TBS (pH7.4) with 1% BSA, 0.02% Proclin300 and 50% Glycerol.

SUBCELLULAR LOCATIONCytoplasmic and Golgi ApparatusSIMILARITYBelongs to the dymeclin family.SUBUNITInteracts with GOLM1 and PPIB.

Post-translational Myristoylated in vitro; myristoylation is not essential for protein targeting to

modifications Golgi compartment.

DISEASE Defects in DYM are the cause of Dyggve-Melchior-Clausen syndrome (DMC)

[MIM:223800]. DMC is a rare autosomal recessive disorder characterized by short trunk dwarfism, microcephaly and psychomotor retardation. Electron microscopic study of cutaneous cells of affected patients shows dilated rough endoplasmic reticulum, enlarged and aberrant vacuoles and numerous vesicles. DMC is progressive. Defects in DYM are the cause of Smith-McCort

dysplasia (SMC) [MIM:607326]. SMC is a rare autosomal recessive osteochondrodysplasia characterized by short limbs and trunk with barrel-shaped chest. The radiographic phenotype includes platyspondyly, generalized abnormalities of the epiphyses and metaphyses, and a distinctive

lacy appearance of the iliac crest, features identical to those of

Dyggve-Melchior-Clausen syndrome.

Important Note This product as supplied is intended for research use only, not for use in

human, therapeutic or diagnostic applications.

Background Descriptions Dyggve-Melchior-Clausen syndrome (DMC), a rare autosomal recessive

disorder, is characterized by microcephaly, short trunk dwarfism and sometime psychomotor retardation. Cutaneous cells of affected individuals show dilated rough endoplasmic reticulum and enlarged vacuoles. The Dyggve-Melchior-Clausen syndrome protein, also designated dymeclin, may play a role in proteoglycan metabolism and intracellular protein digestion. It is a widely expressed multi-pass membrane protein, detected primarily in chondrocytes and fetal brain tissue. Defects in dymeclin are also the cause of Smith-McCort dysplasis syndrome (SMC), which has characteristics identical to those of Dyggve-Melchior-Clausen syndrome.

Additional Information

Gene ID 54808

Other Names Dymeclin, Dyggve-Melchior-Clausen syndrome protein, DYM

Target/Specificity Expressed in most embryo-fetal and adult tissues. Abundant in primary

chondrocytes, osteoblasts, cerebellum, kidney, lung, stomach, heart, pancreas and fetal brain. Very low or no expression in the spleen, thymus, esophagus,

bladder and thyroid gland.

Dilution WB=1:500-2000,IHC-P=1:100-500,IHC-F=1:100-500,ICC=1:100-500,IF=1:100-50

0,ELISA=1:5000-10000

Format 0.01M TBS(pH7.4) with 1% BSA, 0.09% (W/V) sodium azide and 50% Glyce

Storage Store at -20 °C for one year. Avoid repeated freeze/thaw cycles. When

reconstituted in sterile pH 7.4 0.01M PBS or diluent of antibody the antibody

is stable for at least two weeks at 2-4 °C.

Protein Information

Name DYM

Function Necessary for correct organization of Golgi apparatus. Involved in bone

development.

Cellular Location Cytoplasm. Golgi apparatus. Membrane; Lipid-anchor. Note=Sequence

analysis programs clearly predict 1 transmembrane region. However, PubMed:18996921 shows that it is not a stably anchored transmembrane protein but it weakly associates with the Golgi apparatus and shuttles

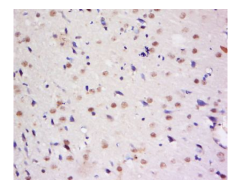
between the Golgi and the cytosol

Tissue Location Expressed in most embryo-fetal and adult tissues. Abundant in primary

chondrocytes, osteoblasts, cerebellum, kidney, lung, stomach, heart, pancreas and fetal brain. Very low or no expression in the spleen, thymus, esophagus,

bladder and thyroid gland

Images



Tissue/cell: Rat brain tissue; 4% Paraformaldehyde-fixed and paraffin-embedded;

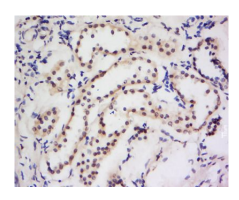
Antigen retrieval: citrate buffer (0.01M, pH 6.0), Boiling bathing for 15min; Block endogenous peroxidase by 3% Hydrogen peroxide for 30min; Blocking buffer (normal goat serum, C-0005) at 37°C for 20 min;

Incubation: Anti- Dymeclin Polyclonal Antibody, Unconjugated(AP55044) 1:200, overnight at 4°C, followed by conjugation to the secondary antibody(SP-0023) and DAB(C-0010) staining

Tissue/cell: Human kidney tissue; 4%

Paraformaldehyde-fixed and paraffin-embedded;

Antigen retrieval: citrate buffer (0.01M, pH 6.0), Boiling



bathing for 15min; Block endogenous peroxidase by 3% Hydrogen peroxide for 30min; Blocking buffer (normal goat serum,C-0005) at 37°C for 20 min; Incubation: Anti- Dymeclin Polyclonal Antibody, Unconjugated(AP55044) 1:200, overnight at 4°C, followed by conjugation to the secondary antibody(SP-0023) and DAB(C-0010) staining

Please note: All products are 'FOR RESEARCH USE ONLY. NOT FOR USE IN DIAGNOSTIC OR THERAPEUTIC PROCEDURES'.