

Product datasheet

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ARG64392 anti-Arylsulfatase B antibody

Package: 100 μg Store at: -20°C

Summary

Product Description Goat Polyclonal antibody recognizes Arylsulfatase B

Tested Reactivity Hu

Tested Application IHC-P, WB

Specificity This antibody is expected to recognise both reported isoforms (NP_000037.2; NP_942002.1).

Host Goat

Clonality Polyclonal

Isotype IgG

Target Name Arylsulfatase B

Species Human

Immunogen C-KLARGHTNGTKPLD

Conjugation Un-conjugated

Alternate Names N-acetylgalactosamine-4-sulfatase; EC 3.1.6.12; MPS6; ASB; G4S; Arylsulfatase B

Application Instructions

Application table	Application	Dilution
	IHC-P	5 - 10 μg/ml
	WB	0.03 - 0.1 μg/ml
Application Note	IHC-P: Antigen Retrieval: Steam tissue section in Citrate buffer (pH 6.0). WB: Recommend incubate at RT for 1h. * The dilutions indicate recommended starting dilutions and the optimal dilutions or concentrations should be determined by the scientist.	

Properties

Form Liquid

Purification Purified from goat serum by antigen affinity chromatography.

Buffer Tris saline (pH 7.3), 0.02% Sodium azide and 0.5% BSA.

Preservative 0.02% Sodium azide

Stabilizer 0.5% BSA

Concentration 0.5 mg/ml

Storage instruction For continuous use, store undiluted antibody at 2-8°C for up to a week. For long-term storage, aliquot

and store at -20°C or below. Storage in frost free freezers is not recommended. Avoid repeated freeze/thaw cycles. Suggest spin the vial prior to opening. The antibody solution should be gently mixed

before use.

Bioinformation

Database links GenelD: 411 Human

Swiss-port # P15848 Human

Background Arylsulfatase B encoded by this gene belongs to the sulfatase family. The arylsulfatase B homodimer

hydrolyzes sulfate groups of N-Acetyl-D-galactosamine, chondriotin sulfate, and dermatan sulfate. The protein is targetted to the lysozyme. Mucopolysaccharidosis type VI is an autosomal recessive lysosomal storage disorder resulting from a deficiency of arylsulfatase B. Two alternatively spliced transcript variants encoding distinct isoforms have been found for this gene. [provided by RefSeq, Jul

2008]

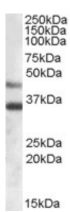
Research Area Cancer antibody; Cell Death antibody; Controls and Markers antibody; Metabolism antibody

Calculated Mw 60 kDa

PTM The conversion to 3-oxoalanine (also known as C-formylglycine, FGly), of a serine or cysteine residue in

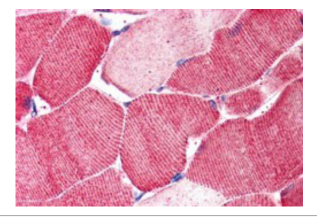
prokaryotes and of a cysteine residue in eukaryotes, is critical for catalytic activity. This post-translational modification is severely defective in multiple sulfatase deficiency (MSD).

Images



ARG64392 anti-Arylsulfatase B antibody WB image

Western Blot: Human Heart lysate (35 μ g protein in RIPA buffer) stained with ARG64392 anti-ARSB antibody at 0.03 μ g/ml dilution.



ARG64392 anti-Arylsulfatase B antibody IHC-P image

Immunohistochemistry: paraffin embedded Human Skeletal Muscle. (Steamed antigen retrieval with citrate buffer pH 6) stained with ARG64392 anti-ARSB antibody at 5 $\mu g/ml$ dilution followed by APstaining.