AMPD1 Polyclonal Antibody

catalog number: E-AB-16156



Note: Centrifuge before opening to ensure complete recovery of vial contents.

Description

Reactivity Human

Immunogen Synthetic peptide of human AMPD1

Host Rabbit **Is otype** IgG

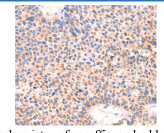
Purification Affinity purification Conjugation Unconjugated

buffer Phosphate buffered solution, pH 7.4, containing 0.05% stabilizer and 50% glycerol.

Applications	Recommended Dilution

WB 1:500-1:2000 IHC 1:100-1:300

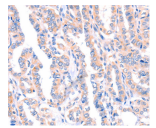
Data



Western Blot analysis of Human fetal muscle tissue using AMPD1 Polyclonal Antibody at dilution of 1:1600

Immunohistochemistry of paraffin-embedded Human liver cancer using AMPD1 Polyclonal Antibody at dilution of 1:70

Calculated-MV:90 kDa



Immunohistochemistry of paraffin-embedded Human thyroid cancer using AMPD1 Polyclonal Antibody at dilution of 1:70

Preparation & Storage

Store at -20°C Valid for 12 months. Avoid freeze / thaw cycles. Storage

Shipping The product is shipped with ice pack, upon receipt, store it immediately at the

temperature recommended.

Background

Adenosine monophosphate deaminase 1 catalyzes the deamination of AMP to IMP in skeletal muscle and plays an important role in the purine nucleotide cycle. Two other genes have been identified, AMPD2 and AMPD3, for the liverand erythocyte-specific isoforms, respectively. Deficiency of the muscle-specific enzyme is apparently a common cause of exercise-induced myopathy and probably the most common cause of metabolic myopathy in the human. Alternatively spliced transcript variants encoding different isoforms have been identified in this gene.

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