

A Reliable Research Partner in Life Science and Medicine

ACTA1 Polyclonal Antibody

catalog number: E-AB-14550

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

Description

Reactivity Human; Mouse; Rat

Immunogen Recombinant protein of human ACTA1

Host Rabbit
Isotype IgG

Purification Affinity purification

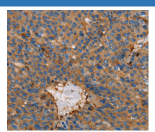
Buffer Phosphate buffered solution, pH 7.4, containing 0.05% stabilizer and 50% glycerol.

Applications Recommended Dilution

WB 1:500-1:2000 **IHC** 1:50-1:200

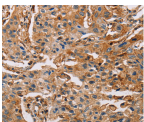
Data





Western Blot analysis of Mouse muscle and heart tissue using Immunohistochemistry of paraffin-embedded Human liver ACTA1 Polyclonal Antibody at dilution of 1:500 cancer using ACTA1 Polyclonal Antibody at dilution of 1:65

Calculated-MW:42 kDa



Immunohistochemistry of paraffin-embedded Human lung cancer using ACTA1 Polyclonal Antibody at dilution of 1:65

Preparation & Storage

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

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

temperature recommended.

Background

For Research Use Only

Elabscience Bionovation Inc.

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The product encoded by this gene belongs to the actin family of proteins, which are highly conserved proteins that play a role in cell motility, structure and integrity. Alpha, beta and gamma actin isoforms have been identified, with alpha actins being a major constituent of the contractile apparatus, while beta and gamma actins are involved in the regulation of cell motility. This actin is an alpha actin that is found in skeletal muscle. Mutations in this gene cause nemaline myopathy type 3, congenital myopathy with excess of thin myofilaments, congenital myopathy with cores, and congenital myopathy with fiber-type disproportion, diseases that lead to muscle fiber defects.

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Rev. V1.7