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## Muscle actin mouse Monoclonal Antibody(3E9)

Catalog NoYP-Ab-02997IsotypeIgGReactivityHuman;Rat;MouseApplicationsWB;IHC;IFGene NameACTA1Protein NameACTA1	
ReactivityHuman;Rat;MouseApplicationsWB;IHC;IFGene NameACTA1	
ApplicationsWB;IHC;IFGene NameACTA1	
Gene Name ACTA1	
Protein Name ACTA1	
Immunogen Synthetic Peptide of Muscle actin	
Specificity Muscle actin protein detects endogenous levels of ACTA1	
Formulation Liquid in PBS containing 50% glycerol, 0.5% BSA and 0.02% sodium azide	e.
Source Monoclonal, Mouse	
PurificationThe antibody was affinity-purified from mouse ascites by affinity-chromatogusing specific immunogen.	graphy
Dilution WB 1:1000-2000, IHC 1:100-200. IF 1:50-200	
Concentration 1 mg/ml	
Purity ≥90%	
Storage Stability -20°C/1 year	
Synonyms ACTA1	
Observed Band 42kD	
Cell Pathway Cytoplasm, cytoskeleton.	
Tissue Specificity Epithelium, Skeletal muscle,	
Function disease:Defects in ACTA1 are a cause of congenital myopathy with excess myofilaments (CM) [MIM:102610].,disease:Defects in ACTA1 are a cause congenital myopathy with fiber-type disproportion (CFTD) [MIM:255310]; a known as congenital fiber-type disproportion myopathy (CFTDM). CFTD is genetically heterogeneous disorder in which there is relative hypotrophy of muscle fibers compared to type 2 fibers on skeletal muscle biopsy. However, these findings are not specific and can be found in many different myopath neuropathic conditions.,disease:Defects in ACTA1 are the cause of nemali myopathy type 3 (NEM3) [MIM:161800]. Nemaline myopathy (NEM) is a for congenital myopathy characterized by abnormal thread- or rod-like structure muscle fibers on histologic examination. The clinical phenotype is highly variable.	of lso type 1 er, nic and ine orm of res in
<b>Background</b> The product encoded by this gene belongs to the actin family of proteins, are highly conserved proteins that play a role in cell motility, structure and integrity. Alpha, beta and gamma actin isoforms have been identified, with	



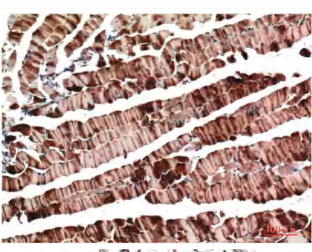
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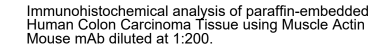


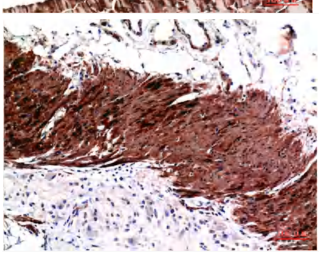
	actins being a major constituent of the contractile apparatus, while beta and gamma actins are involved in the regulation of cell motility. This actin is an alpha actin that is found in skeletal muscle. Mutations in this gene cause nemaline myopathy type 3, congenital myopathy with excess of thin myofilaments, congenital myopathy with cores, and congenital myopathy with fiber-type disproportion, diseases that lead to muscle fiber defects. [provided by RefSeq, Jul 2008],
matters needing attention	Avoid repeated freezing and thawing!
Usage suggestions	This product can be used in immunological reaction related experiments. For more information, please consult technical personnel.

## **Products Images**



Immunohistochemical analysis of paraffin-embedded Mouse Skeletal Muscle Tissue using Muscle Actin Mouse mAb diluted at 1:200.





94KD 66KD 45KD 35KD 26KD 7100 7100 373 20x brain

1

2

3

Western blot analysis of 1) Hela Cell Lysate, 2) 3T3 Cell Lysate, 3) Rat Brain Tissue Lysate using Muscle Actin Mouse mAb diluted at 1:1000.