

OriGene Technologies, Inc.

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Product datasheet for TA803494AM

Lamin A (LMNA) Mouse Monoclonal Antibody (Biotin conjugated) [Clone ID: OTI1B7]

Product data:

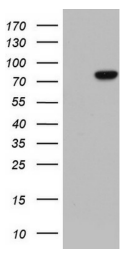
Product Type:	Primary Antibodies
Clone Name:	OTI1B7
Applications:	IF, WB
Recommended Dilution:	WB 1:2000
Reactivity:	Human, Mouse, Rat
Host:	Mouse
lsotype:	lgG1
Clonality:	Monoclonal
Immunogen:	Full length human recombinant protein of human LMNA (NP_733821) produced in HEK293T cell.
Formulation:	PBS (pH 7.3) containing 1% BSA, 50% glycerol and 0.02% sodium azide.
Concentration:	0.5 mg/ml
Purification:	Purified from mouse ascites fluids or tissue culture supernatant by affinity chromatography (protein A/G)
Conjugation:	Biotin
Storage:	Store at -20°C as received.
Stability:	Stable for 12 months from date of receipt.
Predicted Protein Size:	74 kDa
Gene Name:	lamin A/C
Database Link:	<u>NP_733821</u> <u>Entrez Gene 16905 MouseEntrez Gene 60374 RatEntrez Gene 4000 Human</u> <u>P02545</u>



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Background:	The nuclear lamina consists of a two-dimensional matrix of proteins located next to the inner nuclear membrane. The lamin family of proteins make up the matrix and are highly conserved in evolution. During mitosis, the lamina matrix is reversibly disassembled as the lamin proteins are phosphorylated. Lamin proteins are thought to be involved in nuclear stability, chromatin structure and gene expression. Vertebrate lamins consist of two types, A and B. Alternative splicing results in multiple transcript variants. Mutations in this gene lead to several diseases: Emery-Dreifuss muscular dystrophy, familial partial lipodystrophy, limb girdle muscular dystrophy, dilated cardiomyopathy, Charcot-Marie-Tooth disease, and Hutchinson-Gilford progeria syndrome. [provided by RefSeq, Apr 2012]
Synonyms:	CDCD1; CDDC; CMD1A; CMT2B1; EMD2; FPL; FPLD; FPLD2; HGPS; IDC; LDP1; LFP; LGMD1B; LMN1; LMNC
Protein Families	: Druggable Genome
Protein Pathway	rs: Arrhythmogenic right ventricular cardiomyopathy (ARVC), Dilated cardiomyopathy, Hypertrophic cardiomyopathy (HCM)

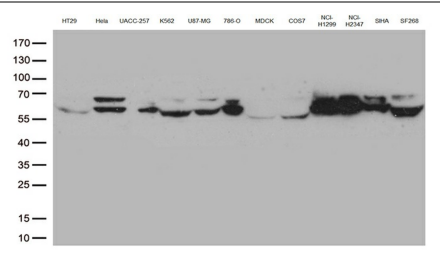
Product images:



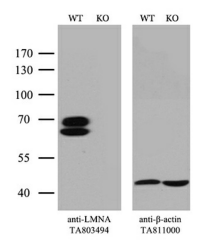
HEK293T cells were transfected with the pCMV6-ENTRY control (Left lane) or pCMV6-ENTRY LMNA ([RC204970], Right lane) cDNA for 48 hrs and lysed. Equivalent amounts of cell lysates (5 ug per lane) were separated by SDS-PAGE and immunoblotted with anti-LMNA. Positive lysates [LY406886] (100ug) and [LC406886] (20ug) can be purchased separately from OriGene.

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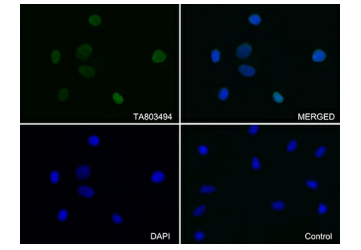
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Western blot analysis of extracts (35ug) from different cell line by using anti-LMNA monoclonal antibody (1:500).



Equivalent amounts of cell lysates (10 ug per lane) of wild-type Hela cells (WT, Cat# LC810HELA) and LMNA-Knockout Hela cells (KO, Cat# [LC810130]) were separated by SDS-PAGE and immunoblotted with anti-LMNA monoclonal antibody [TA803494]. Then the blotted membrane was stripped and reprobed with antib-actin antibody ([TA811000]) as a loading control (1:500).



Immunofluorescent staining of Hela cells using anti-LMNA mouse monoclonal antibody ([TA803494], green, upper left; merged, upper right) or Isotype control (merged, lower right). Cell nuclei were stained with DAPI (blue, lower left) (1:100).

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