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See the following pages for more information!



Lieferung & Zahlungsart

siehe unsere [Liefer- und Versandbedingungen](#)

Zuschläge

- Mindermengenzuschlag
- Trockeneiszuschlag
- Gefahrgutzuschlag
- Expressversand

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LMO1 monoclonal antibody (M06), clone 4C7

Catalog # : H00004004-M06

規格 : [100 ug]

List All

Specification

Product Description: Mouse monoclonal antibody raised against a full length recombinant LMO1.

Immunogen: LMO1 (NP_002306, 15 a.a. ~ 156 a.a) full length recombinant protein with GST tag. MW of the GST tag alone is 26 KDa.

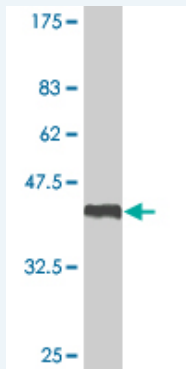
Sequence: VQPKGKQKGCAGCNRKIKDRYLLKALDKYWHEDCLKCACCCDCRLGEVG
STLYTKANLILCRRDYLRFLGTTGNCAACSKLIPAFEMVMRARDNVYHLD
CFACQLCNQRF CVGDKFFLKNMILCQMDYEEGQLNGTFESQVQ

Host: Mouse

Reactivity: Human

Isotype: IgG2a Kappa

Quality Control Testing: Antibody Reactive Against Recombinant Protein.



Western Blot detection against Immunogen (41.62 KDa) .

Storage Buffer: In 1x PBS, pH 7.4

Storage Instruction: Store at -20°C or lower. Aliquot to avoid repeated freezing and thawing.

MSDS: [Download](#)

Datasheet: [Download](#)

Applications

Western Blot (Recombinant protein)

[Protocol Download](#)

ELISA

Gene Information

Entrez GeneID: [4004](#)

Application Image

Western Blot (Recombinant protein)

ELISA

**GeneBank
Accession#:** [NM_002315](#)

**Protein
Accession#:** [NP_002306](#)

Gene Name: LMO1

Gene Alias: MGC116692,RBTN1,RHOM1,TTG1

**Gene
Description:** LIM domain only 1 (rhombotin 1)

Omim ID: [186921](#)

Gene Ontology: [Hyperlink](#)

Gene Summary: LMO1 encodes a cysteine-rich, two LIM domain transcriptional regulator. It is mapped to an area of consistent chromosomal translocation in chromosome 11, disrupting it in T-cell leukemia, although more rarely than the related gene, LMO2 is disrupted. [provided by RefSeq]

**Other
Designations:** LIM domain only 1

Related Disease

[Osteoporosis Tobacco Use Disorder](#)

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