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Research Use Only. Not for diagnostic or therapeutic use.

EB07196-T - Goat Anti-LEF1 Antibody - Trial

Size: 20µg specific antibody in 40µl



Target Protein

Principal Names: LEF1, lymphoid enhancer-binding factor 1, HGNC:6551, DKFZp586H0919, TCF1ALPHA, lymphoid enhancer binding factor-1

Official Symbol: LEF1

Accession Number(s): NP_057353.1; NP_001124185.1; NP_001124186.1;

NP_001159591.1

Human GeneID(s): 51176

Non-Human GenelD(s): 16842 (mouse), 161452 (rat)

Immunogen

Peptide with sequence C-QHEQRKEQEPKRPH, from the internal region of the protein sequence according to NP_057353.1; NP_001124185.1; NP_001124186.1; NP_001159591.1.

Please note the peptide is available for sale.

Purification and Storage

Purified from goat serum by ammonium sulphate precipitation followed by antigen affinity chromatography using the immunizing peptide.

Supplied at 0.5 mg/ml in Tris saline, 0.02% sodium azide, pH7.3 with 0.5% bovine serum albumin

Aliquot and store at -20°C. Minimize freezing and thawing.

Applications Tested

Peptide ELISA: antibody detection limit dilution 1:32000.

Western blot: Approx 50--55kDa band observed in nuclear lysates of cell line K562 (calculated MW of 44.2kDa according to NP_057353.1)/ This molecular weight is routinely observed by other sources and was successfully blocked by incubation with the immunising peptide . Recommended concentration: 1-3μg/ml. . Primary incubation 1 hour at room temperature.

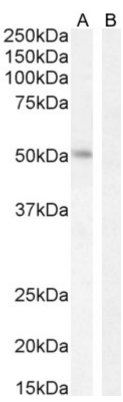
Immunofluorescence: Strong expression of the protein seen in the nuclei of Jurkat and U2OS cells. Recommended concentration: 10µg/ml.

Flow Cytometry: Flow cytometric analysis of Jurkat cells. Recommended concentration: 10ug/ml.

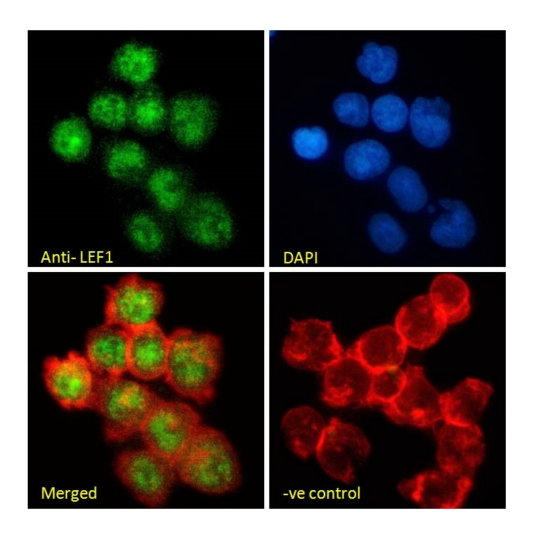
Species Reactivity

Tested: Human

Expected from sequence similarity: Human, Mouse, Rat, Dog, Cow

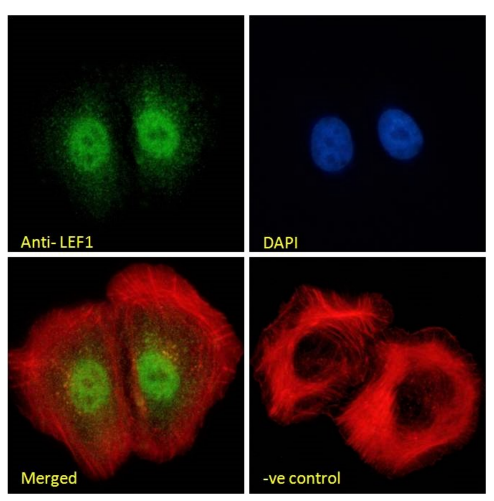


EB07196 ($2\mu g/ml$) staining of K562 nuclear cell lysate (A) + peptide (B). ($35\mu g$ protein in RIPA buffer) Detected by chemiluminescence.

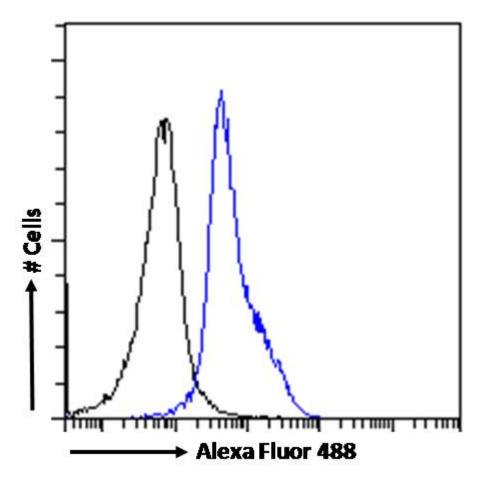


EB07196 Immunofluorescence analysis of paraformaldehyde fixed Jurkat cells, permeabilized with 0.15% Triton. Primary incubation 1hr (10ug/ml) followed by Alexa Fluor 488 secondary antibody (2ug/ml), showing nuclear staining. Actin filaments were stained with phalloidin (red) and the nuclear stain is DAPI (blue). Negative control:

Unimmunized goat IgG (10ug/ml) followed by Alexa Fluor 488 secondary antibody (2ug/ml).



EB07196 Immunofluorescence analysis of paraformaldehyde fixed U2OS cells, permeabilized with 0.15% Triton. Primary incubation 1hr (10ug/ml) followed by Alexa Fluor 488 secondary antibody (2ug/ml), showing nuclear staining. Actin filaments were stained with phalloidin (red) and the nuclear stain is DAPI (blue). Negative control: Unimmunized goat IgG (10ug/ml) followed by Alexa Fluor 488 secondary antibody (2ug/ml).



EB07196 Flow cytometric analysis of paraformaldehyde fixed Jurkat cells (blue line), permeabilized with 0.5% Triton. Primary incubation 1hr (10ug/ml) followed by Alexa Fluor 488 secondary antibody (1ug/ml). IgG control:

Unimmunized goat IgG (black line) followed by Alexa Fluor 488 secondary antibody.