

#### **International Office**

#### **Everest Biotech Ltd**

Vector Laboratories, Inc. 6737 Mowry Ave Newark, CA 94560 United States

**Customer Service:** 

customerservice@vectorlabs.com

Technical Service:

technical@vectorlabs.com

Tel: +1 (800) 227-6666

www.everestbiotech.com

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# EB06097 - Goat Anti-SAE1 / AOS1 Antibody

Size: 100µg specific antibody in 200µl



## **Target Protein**

**Principal Names:** SAE1, AOS1, SUA1, HSPC140, SUMO-1 activating enzyme subunit 1, sentrin/SUMO-activating protein AOS1, SUMO-1 activating enzyme E1 N subunit, ubiquitin-like protein SUMO-1 activating enzyme, SUMO1 activating enzyme subunit 1, FLJ3091, UBLE1A, activator of SUMO1, ubiquitin-like 1-activating enzyme E1A

Official Symbol: SAE1

Accession Number(s): NP\_005491.1

Human GeneID(s): 10055

### **Immunogen**

Peptide with sequence MKGNGIVECLGPK, from the C Terminus of the protein sequence according to NP\_005491.1.

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.

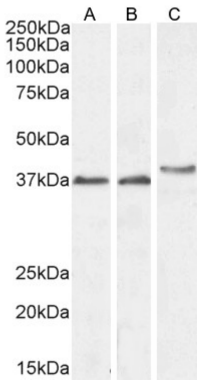
**Western blot:** Approx. 37kDa band observed in lysates of cell lines HEK293 and A549, and approx. 38-40kDa in lysates of cell line Jurkat (calculated MW of 38.4kDa according to NP\_005491.1). Recommended concentration: 0.5-2µg/ml. Primary incubation 1 hour at room temperature.

**Flow Cytometry:** Flow cytometric analysis of A431 cells. Recommended concentration: 10ug/ml.

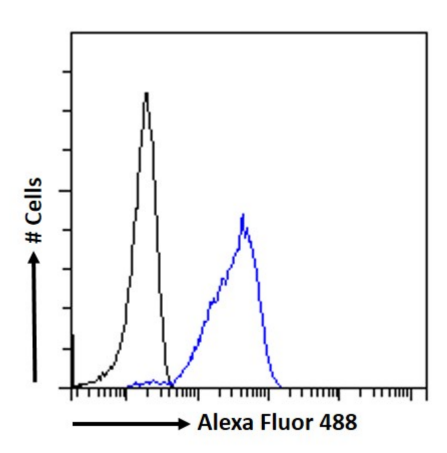
### **Species Reactivity**

Tested: Human

Expected from sequence similarity: Human



EB06097 (1μg/ml) staining of HEK293 (A), A549 (B) and (2ug/ml) Jurkat (C) cell lysate (35μg protein in RIPA buffer). Detected by chemiluminescence.



EB06097 Flow cytometric analysis of paraformaldehyde fixed A431 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.