

# [KO Validated] SMAD2/SMAD3 Rabbit mAb

Catalog No.: A28421

KO Validated

Recombinant

1 Publications

## Basic Information

### Observed MW

52 kDa/60 kDa

### Calculated MW

25-52 kDa

### Category

Primary antibody

### Applications

WB, IP, ChIP, ELISA

### Cross-Reactivity

Human, Mouse, Rat

### CloneNo number

ARC77767

## Background

The protein encoded by this gene belongs to the SMAD, a family of proteins similar to the gene products of the Drosophila gene 'mothers against decapentaplegic' (Mad) and the C. elegans gene Sma. SMAD proteins are signal transducers and transcriptional modulators that mediate multiple signaling pathways. This protein mediates the signal of the transforming growth factor (TGF)-beta, and thus regulates multiple cellular processes, such as cell proliferation, apoptosis, and differentiation. This protein is recruited to the TGF-beta receptors through its interaction with the SMAD anchor for receptor activation (SARA) protein. In response to TGF-beta signal, this protein is phosphorylated by the TGF-beta receptors. The phosphorylation induces the dissociation of this protein with SARA and the association with the family member SMAD4. The association with SMAD4 is important for the translocation of this protein into the nucleus, where it binds to target promoters and forms a transcription repressor complex with other cofactors. This protein can also be phosphorylated by activin type 1 receptor kinase, and mediates the signal from the activin. Alternatively spliced transcript variants have been observed for this gene. The SMAD family of proteins are a group of intracellular signal transducer proteins similar to the gene products of the Drosophila gene 'mothers against decapentaplegic' (Mad) and the C. elegans gene Sma. The SMAD3 protein functions in the transforming growth factor-beta signaling pathway, and transmits signals from the cell surface to the nucleus, regulating gene activity and cell proliferation. This protein forms a complex with other SMAD proteins and binds DNA, functioning both as a transcription factor and tumor suppressor. Mutations in this gene are associated with aneurysms-osteoarthritis syndrome and Loey's-Dietz Syndrome 3.

## Recommended Dilutions

**WB** 1:6000 - 1:15000

**IP** 0.5µg-4µg antibody for  
200µg-400µg extracts of  
whole cells

**ChIP** 2µg antibody for 10µg-15µg  
of Chromatin

**ELISA** Recommended starting  
concentration is 1 µg/mL.  
Please optimize the  
concentration based on your  
specific assay  
requirements. For high-ratio  
antibody dilutions  
(≥1:10000) a sequential  
dilution method is strongly  
recommended to ensure  
measurement accuracy.

## Immunogen Information

### Gene ID

4087/4088

### Swiss Prot

Q15796/P84022

### Immunogen

Synthetic peptide. This information is considered to be commercially sensitive.

### Synonyms

CHTD8; JV18; JV18-1; LDS6; MADH2; MADR2; hMAD-2; hSMAD2; HSPC193; HsT17436; JV15-2; LDS1C; LDS3; MADH3; hMAD-3; hSMAD3; mad3

## Product Information

### Source

Rabbit

### Isotype

IgG

### Purification

Affinity purification

### Storage

Store at -20°C. Avoid freeze / thaw cycles.

Buffer: PBS with 0.09% Sodium azide, 0.05% BSA, 50% glycerol, pH7.3.

Contact

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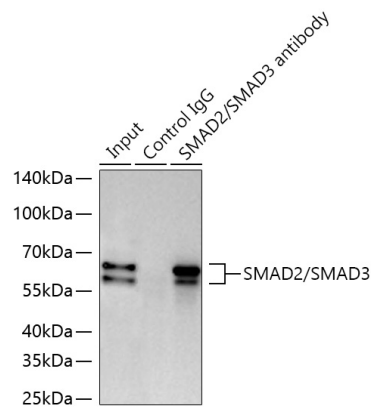
☎ | 400-999-6126

✉ | [cn.market@abclonal.com.cn](mailto:cn.market@abclonal.com.cn)

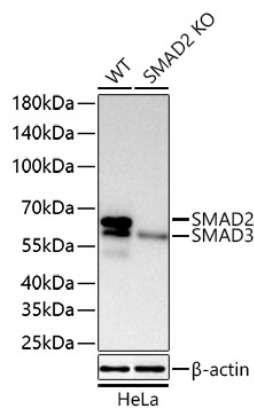
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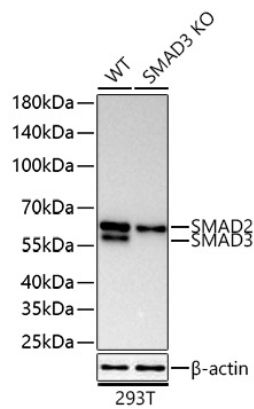
Validation Data



Immunoprecipitation of SMAD2/SMAD3 from 300 µg extracts of NIH/3T3 cells was performed using 0.5 µg of [KO Validated] SMAD2/SMAD3 Rabbit mAb (A28421). Rabbit Control IgG (AC005) was used to precipitate the Control IgG sample. IP samples were eluted with 1x Laemmli Buffer. The Input lane represents 10% of the total input. Western blot analysis of immunoprecipitates was conducted using [KO Validated] SMAD2/SMAD3 Rabbit mAb (A28421) at a dilution of 1:6000.

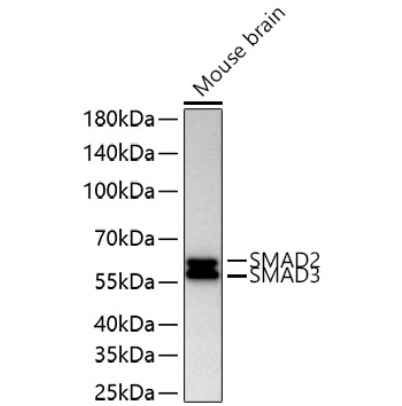


Western blot analysis of lysates from wild type (WT) and SMAD2/SMAD3 knockout (KO) HeLa cells using [KO Validated] SMAD2/SMAD3 Rabbit mAb (A28421) at 1:15000 dilution incubated overnight at 4°C. Secondary antibody: HRP-conjugated Goat anti-Rabbit IgG (H+L) (AS014) at 1:10000 dilution. Lysates/proteins: 25 µg per lane. Blocking buffer: 3% nonfat dry milk in TBST. Detection: ECL Basic Kit (RM00020). Exposure time: 45 s.

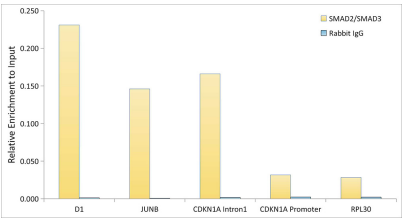


Western blot analysis of lysates from wild type (WT) and SMAD2/SMAD3 knockout (KO) 293T cells using [KO Validated] SMAD2/SMAD3 Rabbit mAb (A28421) at 1:15000 dilution incubated overnight at 4°C. Secondary antibody: HRP-conjugated Goat anti-Rabbit IgG (H+L) (AS014) at 1:10000 dilution. Lysates/proteins: 25 µg per lane. Blocking buffer: 3% nonfat dry milk in TBST. Detection: ECL Basic Kit (RM00020). Exposure time: 45 s.

Validation Data



Western blot analysis of lysates from Mouse brain using [KO Validated] SMAD2/SMAD3 Rabbit mAb (A28421) at 1:15000 dilution incubated overnight at 4°C.  
Secondary antibody: HRP-conjugated Goat anti-Rabbit IgG (H+L) (AS014) at 1:10000 dilution.  
Lysates/proteins: 25 µg per lane.  
Blocking buffer: 3% nonfat dry milk in TBST.  
Detection: ECL Basic Kit (RM00020).  
Exposure time: 45 s.



Chromatin immunoprecipitation was performed with 15 µg of cross-linked chromatin from A549 cells treated with TGF-β1 (20 ng/mL, 30 min), using 2 µg of [KO Validated] SMAD2/SMAD3 Rabbit mAb (A28421) and Rabbit IgG isotype control (AC042). The enrichment of immunoprecipitated DNA at different genomic loci was examined by quantitative PCR. The histogram compares the ratio of the immunoprecipitated DNA to the input at given loci.