

# Affinity Gel-conjugated Mouse anti HA-Tag mAb {Anti-HA □□□□}

Catalog No.: AE059 1 Publications

## atalog No.: AE039 | 1 Publications

## **Basic Information**

#### **Observed MW**

Refer to Figures/60kDa

#### **Calculated MW**

# Category

Tag antibody

# **Applications**

IΡ

### **Cross-Reactivity**

Species independent

#### CloneNo number

AMC0517

# Conjugate

Affinity Gel

# **Background**

Protein tags are peptide sequences genetically grafted onto a recombinant protein. Often these tags are removable by chemical agents or by enzymatic means, such as proteolysis or intein splicing. Tags are attached to proteins for various purposes. Epitope tags are short peptide sequences which are chosen because high-affinity antibodies can be reliably produced in many different species. These are usually derived from viral genes, which explain their high immunoreactivity. Epitope tags include V5-tag, Myc-tag, HA-tag and NE-tag. These tags are particularly useful for western blotting, immunofluorescence and immunoprecipitation experiments, although they also find use in antibody purification.

## **Recommended Dilutions**

Binding Cap 1.5 mg HA protein/mL

**IP** 20μl-40μl Affinity Gel for 100μg-300μg extracts of

whole cells

# Immunogen Information

Gene ID Swiss Prot

#### **Immunogen**

A synthetic peptide corresponding to HA tag.

#### **Synonyms**

HA;HA tag;HA-tag

## **Contact**

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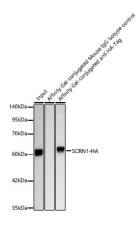
## **Product Information**

SourceIsotypePurificationMouseIgG1,KappaAffinity purification

#### Storage

Store at -20  $^{\circ}\text{C}.$  Avoid freeze / thaw cycles.

Buffer: PBS with 0.02% sodium azide,50% glycerol,pH7.3.



Immunoprecipitation of SCRN1-HA from 300  $\mu g$  extracts of 293T cells transfected with a SCRN1 expression vector containing a single N-terminal HA-Tag was performed using 30  $\mu$ l of Affinity Gelconjugated Mouse anti HA-Tag mAb (AE059). Affinity Gelconjugated Mouse Control IgG pAb 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 Mouse anti HA-Tag mAb (AE065) at a dilution of 1:2000.