

Product Code: sutma

#### **Product Information**

**Description:** The ChromoTek SUMO-Tag-Trap Magnetic Agarose consists of an anti-SUMO-Tag Nanobody/VHH, which is coupled to magnetic agarose beads. It can be used for the immunoprecipitation of SUMO-tagged from cell extracts of various organisms. The ChromoTek SUMO-Tag-Trap can also be used in conjunction with SUMO proteases such as SenP2 for on-bead digestion of SUMO-Tag fusion proteins to release the protein of interest.

Applications: IP, Co-IP

**Specificity/Target:** Binds specifically to all common variants of the SUMO-Tag. The SUMO-Tag is based on Small Ubiquitin-like **Mo**difier (SUMO) proteins of a size of ca. 11 kDa, which are covalently attached to target proteins as a post-translational modification. Fusion of the SUMO-Tag to a protein of interest (POI) may increase expression and solubility of the POI. Also, the SUMO-Tag can be specifically removed by SUMO proteases such as SenP2 without leaving non-native residues behind. At least three SUMO variants are commonly used as SUMO-Tag and are all recognized by the ChromoTek SUMO-Tag-Trap: the yeast SUMO homolog SMT3, the human SUMO3 and SUMOStar, a version of SMT3 resistant to SUMO proteases. Please note that the ChromoTek SUMO-Tag-Trap will also bind non-discriminatorily to endogenous SUMO homologs such as SUMO1, SUMO2 and SUMO3 present in human cells.

Binding capacity: . 30 µg of recombinant SUMO-tagged protein (~40 kDa) per 25 µL bead slurry

Bead Size: 40 µm (cross-linked 6 % magnetic agarose beads)

Elution Buffer: 2x SDS-sample buffer (Lämmli), 200 mM glycine pH 2.5

Wash Buffer Compatibility: 1 M NaCl, 5 mM DTT, 5 mM β-mercaptoethanol, 5 mM TCEP, 2% NP40,

2% Triton X-100, 0.0% SDS, 2 M Urea

**Type:** Nanobody

Class: Recombinant

**Host:** Alpaca

**Shipment:** Shipped at ambient temperature

Storage Buffer: 20 % ethanol

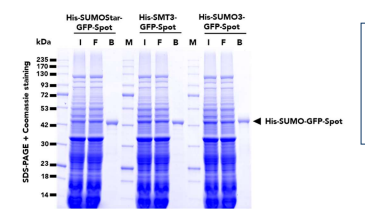
**Storage Condition:** Upon receipt store at +4°C. Do not freeze!

**Stability:** Stable for 1 year upon receipt



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#### **Selected Validation Data**



Immunoprecipitation of three different variants of SUMO-tagged, GFP fusion proteins E. coli cell lysates using SUMO-Tag-Trap Magnetic Agarose. I: Input, F: Flow-through, B: Bound.

## **Suggested Buffer Compositions for IP**

Buffer	Composition	
Lysis Buffer	10 mM Tris/Cl pH 7.5, 150 mM NaCl, 0.5 mM EDTA, 0.5 %	
	Nonidet™ P40 Substitute (adjust the pH at +4°C)	
RIPA Buffer	10 mM Tris/Cl pH 7.5, 150 mM NaCl, 0.5 mM EDTA, 0.1 % SDS, 1	
	% Triton™ X-100, 1 %	
	deoxycholate (adjust the pH at +4°C)	
Dilution Buffer	10 mM Tris/Cl pH 7.5, 150 mM NaCl, 0.5 mM EDTA (adjust the pH	
	at +4°C)	
Wash Buffer	10 mM Tris/Cl pH 7.5, 150 mM NaCl, 0.05 % Nonidet™ P40	
	Substitute, 0.5 mM EDTA (adjust the pH at +4°C)	
2x SDS-sample buffer	120 mM Tris/Cl pH 6.8, 20 % glycerol, 4 % SDS, 0.04 %	
	bromophenol blue, 10 % β- mercaptoethanol	
Acidic elution buffer	200 mM glycine pH 2.5 (adjust the pH at +4°C)	
Neutralization buffer	on buffer 1 M Tris pH 10.4 (adjust the pH at +4°C)	

Note: Use your equivalent cell lysis buffer for other cell types like yeast, plants, insects, bacteria. Consider using a Wash buffer without detergent for Co-IP.

#### **Product Sizes**

Product	Product Code	Size
SUMO-Tag-Trap Magnetic Agarose	sutma-10	10 reactions
	sutma-20	20 reactions
	sutma-100	100 reactions
	sutma-200	200 reactions
	sutma-400	400 reactions



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### Protocol at a glance

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- Perform all steps at 4°C
- Use your preferred cell lysis buffer and cell lysis conditions

#### **Cell Lysis**

General



- Use 10<sup>6</sup>-10<sup>7</sup> cells and 200 μL Lysis buffer.
- · Perform cell lysis and clear lysate
- Mix 200 μl cleared lysate with 300 μL dilution buffer.

### Bead Equilibration



- Transfer 25 µL bead slurry into a 1.5 mL tube
- Equilibrate beads 3x with 500 μL dilution buffer

### Protein binding



- Add 500 µL diluted lysate to beads
- Rotate end-over-end for 1 hour at 4°C.

#### Washing



- Wash beads 3x with 500 µL wash buffer
- Transfer beads to a new tube during the last washing step

### Elution with SDS-sample buffer



- Resuspend beads in 80 µL 2x SDSsample buffer
- Boil beads for 5 min at 95°C
- Analyze the supernatant in SDS-PAGE/ Western Blot



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### **Immunoprecipitation Protocol**

#### **Cell Material**

The following protocol describes the preparation of a mammalian cell lysate.

For other type of cells, we recommend using 500  $\mu g$  of cell extract and start the protocol with step Bead equilibration.

#### **Mammalian Cell Lysis**

**Note:** Harvesting of cells and cell lysis should be performed with ice-cold buffers. We strongly recommend adding protease inhibitors to the Lysis buffer to prevent degradation of your target protein and its binding partners.

For one immunoprecipitation reaction, we recommend using  $\sim 10^6$ - $10^7$  cells.

- 1. Choice of lysis buffer:
- a. For cytoplasmic proteins, resuspend the cell pellet in 200  $\mu$ L ice-cold Lysis buffer by pipetting up and down. Supplement Lysis buffer with protease inhibitor cocktail and 1 mM PMSF (not included).
- b. For nuclear/chromatin proteins, resuspend cell pellet in 200  $\mu$ L ice-cold RIPA buffer supplemented with DNase I (f.c. 75-150 Kunitz U/mL), MgCl<sub>2</sub> (f.c. 2.5 mM), protease inhibitor cocktail and PMSF (f.c. 1 mM) (not included).
- 2. Place the tube on ice for 30 min and extensively pipette the suspension every 10 min.
- 3. Centrifuge cell lysate at 17,000x g for 10 min at  $+4^{\circ}$ C. Transfer cleared lysate (supernatant) to a pre-cooled tube and add 300  $\mu$ L Dilution buffer supplemented with 1 mM PMSF and protease inhibitor cocktail (not included). If required, save 50  $\mu$ L of diluted lysate for further analysis (input fraction).

#### **Bead Equilibration**

- 1. Resuspend the beads by gently pipetting them up and down or by inverting the tube. Do not vortex the beads!
- 2. Transfer 25  $\mu$ L of bead slurry into a 1.5 mL reaction tube.
- 3. Add 500 µL ice-cold Dilution buffer.
- 4. Separate the beads with a magnet until the supernatant is clear. Discard the supernatant.

#### **Protein Binding**

- 1. Add diluted lysate to the equilibrated beads.
- 2. Rotate end-over-end for 1 hour at  $+4^{\circ}$ C.

#### Washing

- 1. Separate the beads with a magnet until the supernatant is clear.
- 2. If required, save 50 µL of supernatant for further analysis (flow-through/non-bound fraction).
- 3. Discard remaining supernatant.
- 4. Resuspend beads in 500 μL Wash buffer.



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- 5. Separate the beads with a magnet until the supernatant is clear. Discard the remaining supernatant.
- 6. Repeat this step at least twice.
- 7. During the last washing step, transfer the beads to a new tube.

Optional: To increase stringency of the Wash buffer, test various salt concentrations e.g. 150-500 mM, and/or add a non-ionic detergent e.g. Triton™ X-100 (see Wash buffer compatibility information for maximal concentrations).

#### **Elution with 2x SDS-sample buffer (Laemmli)**

- 1. Remove the remaining supernatant.
- 2. Resuspend beads in 80 μL 2x SDS-sample buffer.
- 3. Boil beads for 5 min at +95°C to dissociate immunocomplexes from beads.
- 4. Separate the beads with a magnet.
- 5. Analyze the supernatant in SDS-PAGE / Western Blot.

#### **Elution with Acidic Elution Buffer**

- 1. Remove the remaining supernatant.
- 2. Add 50-100  $\mu$ L Acidic elution buffer and constantly pipette up and down for 30-60 sec at +4°C or room temperature.
- 3. Separate the beads with a magnet until the supernatant is clear.
- 4. Transfer the supernatant to a new tube.
- 5. Immediately neutralize the eluate fraction with 5-10 μL Neutralization buffer.
- 6. Repeat this step at least once to increase elution efficiency.

**Note:** Elution at room temperature is more efficient than elution at +4°C. Prewarm buffers for elution at room temperature.

#### **Related Products**

Product	Code
SUMO-Tag-Trap Agarose	suta
SUMO-Tag-Trap Agarose Kit	sutak
SUMO-Tag-Trap Magnetic Agarose Kit	sutmak



# chromotek<sup>®</sup> SUMO-Tag-Trap Magnetic Agarose

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