

PRODUCT INFORMATION

Mouse anti-GST tag mAb Sepharose Purification kit

For mammalian expression system

v. 230201

Catalog number	С07004-К01 / С07004-К02
Package	5 rxns / 10 rxns
Description	Affinity purified anti-GST antibodies, developed in mouse, were conjugated to NHS-sepharose. It is an efficient technique for isolating recombinant proteins or other proteins. The GST epitope system relies on a total 211 amino acids long recombinant antibody, which is able to react with N- and C- terminal GST-tagged fusion proteins and may be used for the immunoprecipitation or immune affinity purification. The purified antibody is immobilized at 3-5 mg antibody per mL of 50% slurry and this kit allows a rapid and efficient affinity purification of active GST fusion proteins. The affinity resin allows an efficient binding of GST fusion proteins without the need for preliminary steps and calibrations. The affinity bound GST fusion proteins can be used for characteristic analysis.

Component	Reagents & Materials	Quantity for 10 rxns (C07004- K02)	Quantity for 5 rxns (C07004- K01)	composition
	Mouse anti-GST	2 mL X 1	1 mL X 1	50% slurry of Mouse anti-
	tag mAb	vial	vial	GST tag mAb sepharose in
	sepharose			1X Phosphate Buffered Saline
	Wash Buffer (10X	5.0 mL X 1	2.5 mL X 1	100 mM Tris/CI pH 7.5;
	concentration)	vial	vial	1.5 M NaCl;
				5 mM EDTA
	Elution Buffer	10 mL X 1 vial	5 mL X 1 vial	0.1 M Glycine pH 2.7
	Neutralization Buffer	2 mL X 1 vial	1 mL X 1 vial	2 M Tris pH 8.0
	spin column	10 pcs	5 pcs	
	collection tube	10 tubes	5 tubes	
	Note: Sepharose is containing 0.05% soo	s 1:1 suspens dium azide as a	ion in Phospl a preservative.	nate Buffered Saline, pH 7.4,
Product capacity	The binding and elur are commonly more buffers for optimal re	tion capacity c than 1 mg of sults is recomr	of 1 mL Mouse GST fusion p nended.	e anti-GST tag mAb sepharose roteins. Trying different elution

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	Micro-centrifuge capable of 15,000 x g			
	1.5 mL Centrifuge tubes			
Materials Required	End-over-end rotator			
but Not Provided	• CoIP Lysis Buffer (mild reaction): 10 mM Tris/CI pH 7.5; 150 mM NaCI; 0.5			
	mM EDTA; 0.5% NP-40			
	• RIPA (vigorous reaction): 100 mM Tris/CI pH 7.5; 300 mM NaCI; 0.2%			
	Sodium Deoxycholate (or 0.1% SDS); 2% NP-40			
Storage	For sustainable use and long term storage, store at 2 °C to 8 °C. DO NOT FREEZE.			
Precautions and Disclaimer	This product is for R&D use only, not for drug, household, or other uses.			
	General notes			
	The Mouse anti-GST tag mAb sepharose is stored in Phosphate Buffered Saline			
	containing 0.05% sodium azide. The PBS must be removed before use and the			
	resin should be equilibrated with 1X Wash Buffer. The equilibration can be			
	performed at room temperature or at 2-8 °C. The Wash Buffer is original stock			
	concentration. Dilute Wash Buffer (10X concentration) to 1X working			
	concentration with distilled water.			
	In the case of bulk reaction. Users can make a pre-reaction through mixing			
	sample and sepharose in 15 mL / 50 mL tube, and then transfer the mixture into			
	the column for binding.			
Procedure	Suggestions on purification of GST fusion proteins			
	1. Cellular debris and particulate matter must be removed by centrifugation or filtration prior to purification on the column.			
	2. Highly viscous samples which may containing chromosomal DNA or RNA			
	should be sonicated or treated with nuclease to decrease the viscosity.			
	3. Perform all steps on ice.			
	A. Sample preparation (Lysis of Mammalian Cells)			
	1. Detach the cells from the culture dish and collect the cell suspension into the centrifuge tube.			
	2. Centrifuge the cell suspension at 400 x g for 5 minutes to pellet the cells.			
	Carefully remove and discard the supernatant.			
	3. Wash cells by re-suspending the cell pellet in ice-cold PBS.			



- Centrifuge the cell suspension at 400 x g for 5 minutes to pellet the cells.
 Carefully remove and discard the supernatant.
- 5. Add 200 μ L of Lysis Buffer to the cell pellet and vortex.
- 6. Incubate the sample for 15 minutes on ice.
- 7. Remove cell debris by centrifugation at 15,000 x g for 5 minutes at 4°C.
- B. Column preparation
- 1. Place an empty spin column on the collection tube.
- 2. Wash the column with 200 μL Wash Buffer.
- 3. Allow the buffer to drain from the column and leave residual Wash Buffer in the column to aid in packing the Mouse anti-GST tag mAb sepharose, then discard the buffer in the collection tube.
- C. Packing the Column

Procedure

- 1. Completely suspend the vial of Mouse anti-GST tag mAb sepharose.
- 2. Transfer 200 μ L volume to an empty centrifuge tube, and wash the sepharose with 1 mL Wash Buffer.
- Spin down the sepharose with 100 x g, 30 seconds' centrifugation and discard supernatant.
 - 4. Immediately transfer the sepharose to the spin column. Allow the sepharose bed to settle. Please prevent the sepharose bed from getting dried.(Note: Make sure the column filter is fixed in the correct position before transferring the sepharose).
 - D. Binding GST fusion protein to the column
 - 1. Dilute the sample with Wash Buffer in 1:3 proportion.
 - Load the sample on spin the column and centrifuge the column at 100 x g for 30 seconds. Users can also perform this binding reaction in a new 1.5 mL centrifuge tube.

(Note: Depending upon the GST fusion protein and the flow rate, not all of the protein may bind. Repeat loading the sample to increase binding efficiency).

- 3. Collect the fractions using empty centrifuge tube.
- 4. Wash the spin column with 300 μL Wash Buffer more than 6 times.

(Note: To eliminate the noisy band in sample, more washing step is recommended).

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- E. Elution of GST fusion protein
- 1. Add 5 x 100 μ L Elution Buffer to elute the bound GST fusion protein from the spin column to the collection tube. This step can be supported by a centrifugation at 100 x g for 30 seconds.
- Immediately neutralize the eluted sample by adding 10 µL Neutralization Buffer. Assay sample concentration by measuring the absorbance at 280 nm and combine the fractions with highest absorbance.

(Note: Measuring the absorbance after each Elution move can help collecting the sample more accurately).

F. Optional instead of elution step

Resuspended Mouse anti-GST tag mAb sepharose in 100 μ L 2 x SDS-Sample Buffer for 10 minutes at 95°C to dissociate immune-complexes from Mouse anti-GST tag mAb sepharose. Mouse anti-GST tag mAb sepharose can be collected by centrifugation at 2500 x g for 2 minutes at 4°C and SDS-PAGE is performed with the supernatant.

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