

# Protein A HP MultiTrap

## Instructions for Use

#### **Protein A HP MultiTrap contains**

- 4 prepacked Protein A HP MultiTrap™ 96-well filter plates
- Instructions for use

#### Introduction

Protein A HP MultiTrap is designed for small-scale sample preparation for single use, for example upstream of gel electrophoresis, liquid chromatography and mass spectrometry. MultiTrap may be used with robotic systems or manually, with centrifugation or vacuum.

The plate wells contain Protein A Sepharose™ High Performance columns. The plates are designed for two different applications:

- Enrichment of target proteins
- Purification of antibodies

The current instructions provide background information, protocols, and general useful information for both applications.

#### General handling of the MultiTrap plate

**Centrifugation/vacuum:** Centrifuge the MultiTrap plates or use vacuum. If vacuum is used, apply 0.15 bar until the wells are empty, then slowly increase the vacuum to -0.3 bar (do not apply more vacuum than -0.5 bar). Turn off the vacuum after approximately 5

**Medium:** Mix briefly before removal of liquid in the equilibration, wash and elution steps to increase the efficiency of the step. Incubating on a plate shaker is recommended.

**Incubations:** During incubation, cover the plate using a sealing tape or an appropriate 96-well cover.

**Collection plates:** Collection plates are not included and must be ordered separately (see *Ordering information, on page 5*). Remember to change or empty the collection plate between steps.

**Sample pretreatment:** Excessive cellular debris and lipids may clog the column. Clarify the sample by centrifugation or filtration before applying to the MultiTrap plate MultiTrap plate well.

**Sample pre-treatment:** To prevent target protein degradation, inhibition of protease activity may be required (a Protease Inhibitor Mix is available, see *Ordering information*, on page 5.

#### **Antibody purification**

#### **Purpose**

The Protein A HP MultiTrap 96-well plates are designed for rapid small-scale antibody purification of multiple samples in parallel, for example in antibody screening experiments.

#### **Principle**

Protein A Sepharose HP has a high protein binding capacity and is compatible with all buffers commonly used in antibody purification. The MultiTrap can be used with a standard centrifuge and one purification takes less than 20 minutes. Cell culture supernatants, as well as serum samples, may be directly applied to the wells without prior clarification.

#### Advice on handling

#### **Optimization of parameters**

The parameters for antibody purification may require optimization. Examples of parameters which may require optimization are:

- sample pretreatment
- amount of antibody to be purified
- incubation time
- choice of buffers
- number of washes

#### Sample pretreatment

Antibodies from several species can be purified with Protein A Sepharose High Performance.

IgG from many species has a medium to strong affinity for Protein A at approximately pH 7.0, see *Antibody binding to Protein A and protein G, on page 2*.

The sample should have a pH around 7 before applying to a well. It is therefore important to check the pH of the sample, and adjust it as necessary before applying the sample to the well.

#### **Choice of buffers**

The following buffers are recommended.

Binding buffer: 20 mM sodium phosphate, pH 7.0

Elution buffer: 0.1 M glycine-HCl, pH 2.7

Neutralizing buffer: 1 M Tris-HCl, pH 9.0

**Note:** Use high-purity water and chemicals for buffer preparation.

- Recommended buffers can be easily prepared using Ab Buffer Kit, see Ordering information, on page 5.
- Protein A Sepharose High Performance binds IgG over a wide pH range with a strong affinity at neutral pH. To elute the IgG, it is necessary to lower the pH to about 2.5 to 3.0 depending on the antibody.

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As a safety measure to preserve the activity of acid-labile lgGs, we recommend the addition of 1 M Tris-HCl, pH 9.0, to collection plate used for collecting lgG-containing fractions (60 to 200 μL/ml eluted fraction). In this way, the final pH of the sample will be approximately neutral.

#### **Antibody recovery**

- If the pH of the sample is too low the antibody may have low binding to Protein A Sepharose High Performance matrix. Ensure that the pH is approximately 7.
- If the MultiTrap wells does not have enough capacity for the amount of antibody in the sample the recovery will be less than expected. Decrease the amount of sample added to each well.

# Antibody binding to Protein A and protein G Relative binding strengths for protein A and protein G

Species	Subclass	Protein A binding	Protein G binding
Human	IgA	variable	-
	IgD	-	-
	IgD	-	-
	IgG <sub>1</sub>	++++	++++
	$IgG_2$	++++	++++
	IgG <sub>3</sub>	-	++++
	IgG₄	++++	++++
	IgM	variable	-
Avian egg yolk	IgY	-	-
Cow		++	+++
Dog		++	+
Goat		-	++
Guinea pig	IgG <sub>1</sub>	++++	++
	$IgG_2$	++++	++
Hamster		+	++
Horse		++	++++
Koala		-	+
Llama		-	+
Monkey		++++	++++
(rhesus)			
Mouse	IgG <sub>1</sub>	+	++++
	$IgG_{2a}$	++++	++++
	IgG <sub>2b</sub>	+++	+++
	IgG <sub>3</sub>	++	+++
	IgM	variable	-
Pig		+++	+++
Rabbit		++++	+++
Rat	IgG <sub>1</sub>	-	+
	IgG <sub>2a</sub>	-	++++
	IgG <sub>2b</sub>	-	++
	$IgG_3$	-	++
Sheep		+/-	++

++++ = strong binding

++ = medium binding

- = weak or no binding

#### **Antibody purification protocol**

The protocol may need optimization for your application, see *Advice* on handling, on page 1. Please refer to *General handling of the MultiTrap plate, on page 1* for general handling instructions.

#### Step Action

#### Prepare collection plates

For step 6, prepare 2 collection plates for eluted fractions, each containing 15  $\mu$ L neutralizing buffer per well.

#### 2 Remove storage solution

- Suspend the medium by gently shaking the plate upside down.
- **b.** Remove top and bottom seals and place the MultiTrap plate on a collection plate.
- **c.** Remove the storage solution by centrifugation for 1 min at  $70-100 \times g$ .

#### 3 Equilibrate

- a. Add 300 µL binding buffer and mix briefly.
- **b.** Centrifuge for 30 s at 70-100 × g.

#### 4 Bind antibody

- a. Add maximum 300 µL of the antibody solution.
- b. Incubate for 4 min while gently mixing.
- c. Centrifuge for 30 s at 70-100 × g.

#### Note:

Several sample applications can be made subsequently as long as the capacity of the column is not exceeded.

#### 5 Wash

- a. Add 300  $\mu$ L binding buffer, mix briefly and centrifuge for 30 s at 70-100 × g.
- **b.** Perform this step 2 times total.

#### 6 Elute antibody

- a. Replace the collection plate with a collection plate prepared in step 1.
- **b.** Add 200  $\mu$ L of elution buffer, mix briefly and centrifuge for 30 s at 70 × g and collect the eluate.
- c. Perform this procedure 2 times total.

#### Note:

Most of the bound antibody is eluted after two elution steps.

#### **Protein enrichment**

#### **Purpose**

The Protein A HP MultiTrap prepacked 96-well plates are designed for small-scale protein enrichment for single use, for example for use upstream of gel electrophoresis, liquid chromatography, and mass spectrometry.

#### **Principle**

There are two protocols for protein enrichment using Protein A HP MultiTrap prepacked 96-well plates:

#### **Cross-link protocol**

In the cross-link protocol the protein capturing antibodies are covalently bound to the Protein G Sepharose High Performance matrix by using a cross-linking agent.

The protein of interest is enriched from the sample, purified through washings, and eluted from the wells whereas the antibody remains bound to the matrix.

Use the cross-link protocol:

- If the desired protein/antigen has similar molecular weight as the heavy or light chain of the antibody, which causes problem with comigration in SDS-PAGE analysis.
- If the antibody interferes with downstream analysis.

#### Classic protocol

In the classic protocol protein capturing antibodies are immobilized by binding to Protein A in the Protein A Sepharose High Performance matrix. The classic protocol requires that the capturing antibody used binds to Protein A.

The protein/antigen of interest is enriched from the sample, purified through washings and eluted from the wells together with the antibody.

#### Advice on handling

#### **Optimization of parameters**

The optimal parameters for protein enrichment are dependent on the specific antibody-antigen combination. Optimization may be required for each specific antibody-antigen combination to obtain the best results.

Examples of parameters which may require optimization are:

- Sample pre-treatment
- Amount of protein (antigen) to be enriched
- Incubation time
- Choice of buffers
- Number of washes

#### Sample pretreatment

- Excessive cellular debris and lipids may clog the well. Clarify the sample by centrifugation or filtration before applying to the MultiTrap wells.
- To prevent target protein degradation, inhibition of protease activity may be required (a Protease Inhibitor Mix is available, see Ordering information, on page 5.

#### **Incubation time**

At room temperature, the reaction is usually completed within 30 to 60 min. If the binding is performed at 4°C, it can be left overnight.

#### **Choice of buffers**

It is recommended to use the listed buffers for the indicated type of protocol. A Protein A/G Buffer Kit is available as an accessory for increased convenience. If optimization is required try to use the alternative buffers.

#### Cross-link protocol

Binding buffer: Wash buffer: Elution buffer: Cross-link solutions: TBS (50 mM Tris, 150 mM NaCl, pH 7.5) TBS with 2 M urea, pH 7.5

0.1 M glycine with 2 M urea, pH 2.9

- 200 mM triethanolamine, pH 8.9
- 50 mM DMP (Dimethyl pimelimidate dihydrochloride) in 200 mM triethanolamine, pH 8.9
- 100 mM ethanolamine, pH 8.9

Classic protocol

Binding buffer: Wash buffer:

TBS (50 mM Tris, 150 mM NaCl, pH 7.5)

TBS

Elution buffer: 2.5% acetic acid

**Alternative buffers** 

Wash buffer:

Elution buffer:

- TBS (mild wash)
- TBS with 1% octylglucoside, pH 7.5
- 0.1 M triethanolamine, 0.5 M NaCl, pH 9.0

0.1 M glycine, pH 2.5 to 3.1

- 0.1 M citric acid, pH 2.5 to 3.1
- 2% SDS
- 0.1 M ammonium hydroxide, pH 10 to 11

#### Protein recovery and specific purity

- Improve the specific purity by adding detergent, different salts, and different concentrations of salts to the wash buffer.
- Avoid acidic elution conditions since this may cause low protein vield.
- Minimize impurities that may co-elute with the target protein by adding a preclearing step before the enrichment procedure. For preclearing, use a MultiTrap well that has not been coupled with an antibody. Add the sample and incubate for 0.5 to 4 h. Collect the sample by centrifugation and proceed with the standard protocol using the coupled medium.
- Try alternative buffers, see Section Choice of buffers, on page 3.

#### Additional options when using the classic protocol

 Incubate the antibody with the sample to form an antibodyantigen complex before applying the sample to the well.
 The complex is then applied to the well for binding.

#### **Cross-link protocol**

The protocol may need optimization for your application, see *Advice* on handling, on page 3. Please refer to *General handling of the MultiTrap plate, on page 1* for general handling instructions.

#### Step Action

#### 1 Remove storage solution

- a. Suspend the medium by gently shaking the plate upside down
- **b.** Remove top and bottom seals and place on a collection plate.
- **c.** Remove the storage solution by centrifugation for 1 min at  $700 \times g$ .

#### 2 **Equilibrate**

- a. Add 400 µL binding buffer, mix briefly and centrifuge for 1 min at 700 × g to equilibrate the medium.
- b. Perform this step 3 times total.

#### Step Action

#### 3 Bind antibody

- a. Immediately after equilibration, add 200 µL of the antibody solution per well (0.5 to 1.0 mg/mlLin binding buffer).
- b. Incubate on shaker for 30 min.
- **c.** Centrifuge for 1 min at  $700 \times g$  to remove unbound antibody.

#### 4 Wash

- a. Add 400 µL binding buffer and mix briefly.
- **b.** Centrifuge for 1 min at 700 × g.

#### 5 Change buffer

- a. Add 400 µL triethanolamine and mix briefly.
- **b.** Centrifuge for 1 min at 700 × g.

#### 6 Cross-link

- a. Add 400 µL DMP in triethanolamine.
- **b.** Incubate on shaker for 30 min for 60 min.
- c. Centrifuge for 1 min at 700 × g.

#### 7 Wash

- a. Add  $400 \, \mu L$  triethanolamine and mix briefly.
- **b.** Centrifuge for 1 min at 700 × g.

#### 8 Block

- a. Add 400 µL ethanolamine
- b. Incubate on shaker for 15 min.
- c. Centrifuge for 1 min at 700 x g.

#### 9 Remove unbound antibody

- a. Add 400  $\mu L$  elution buffer and mix briefly.
- **b.** Centrifuge for 1 min at 700 × g.

#### 10 Wash

- a. Add 400  $\mu$ L binding buffer, mix briefly and centrifuge for 1 min at 700 × g.
- **b.** Perform this step 2 times total.

#### 11 Bind target protein

- a. Add 200  $\mu L$  of sample in binding buffer.
- **b.** Incubate on shaker for 60 min.
- c. Replace the collection plate with a clean collection plate
- **d.** Centrifuge for 1 min at 700 × g to collection out unbound sample.
- e. During optimization/trouble shooting: Collect flowthrough.

#### 12 Wash

- $\textbf{a.} \ \ \mathsf{Replace} \ \mathsf{the} \ \mathsf{collection} \ \mathsf{plate} \ \mathsf{with} \ \mathsf{a} \ \mathsf{clean} \ \mathsf{collection} \ \mathsf{plate}$
- **b.** Add  $400 \,\mu\text{L}$  collection buffer, mix briefly and centrifuge for 1 min at  $700 \times g$ .
- c. Perform this step 5 times total.

#### 13 Elute

a. Collect the eluates in separate collection plates.

#### Step Action

- **b.** Add  $200 \,\mu\text{L}$  of elution buffer, mix briefly and centrifuge for 1 min at  $1000 \times g$ .
- c. Perform this step three 3 times total.

#### Classic protocol

The protocol may need optimization for your application, see *Advice* on handling, on page 3. Please refer to *General handling of the MultiTrap plate, on page 1* for general handling instructions.

#### Step Action

#### Remove storage solution

- Suspend the medium by gently shaking the plate upside down.
- **b.** Remove top and bottom seals and place the MultiTrap plate on a collection plate.
- **c.** Remove the storage solution by centrifugation for 1 min at  $700 \times g$ .

#### 2 Equilibrate

- a. Add  $400\,\mu\text{L}$  binding buffer per well, mix briefly and centrifuge for 1 min at  $700\times g$  to equilibrate the medium.
- b. Perform this step 3 times total.

#### 3 Bind antibody

- a. Immediately after equilibration, add 200 µL of the antibody solution per well (0.5 to 1.0 mg/ml in binding buffer).
- **b.** Incubate on shaker for 30 min.
- Centrifuge for 1 min at 700 × g to remove unbound antibody.

#### 4 Wash

- a. Add 400 µL binding buffer per well, mix briefly.
- **b.** Centrifuge for 1 min at 700 × g.

#### 5 Bind target protein

- a. Add 200 µL sample in binding buffer per well.
- **b.** Incubate on shaker for 60 min.
- c. Replace the collection plate with a clean collection plate
- **d.** Centrifuge for 1 min at 700 × g to wash out unbound sample.
- e. During optimization/trouble shooting: Collect flowthrough.

#### Wasi

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- a. Replace the collection plate with a clean collection plate.
   Collect and save washes in case troubleshooting is needed.
- **b.** Add  $400 \, \mu L$  wash buffer per well, mix briefly and centrifuge for 1 min at  $700 \times g$ .
- **c.** Perform this step 5 times total.

Step	Ac	tion
7	Eli	ute
	a.	Collect the eluates in separate collection plates.
	b.	Add 200 $\mu L$ of desired elution buffer per well and shake for 1 min.
	c.	Centrifuge for 1 min at 700 x g.
	d.	Perform this procedure 3 times total.

#### **Characteristics**

Matrix	Highly cross-linked agarose, 6%
Medium	Protein A Sepharose High Performance
Ligand	Native protein A
Ligand coupling method	N-hydroxysuccinimide activation
Ligand density	approx. 3 mg protein A/ml medium
Binding capacity <sup>1</sup>	approx. 20 mg human IgG/ml medium
Average particle size	34 µm
pH stability <sup>2</sup>	3 to 9 (long term)
	2 to 9 (short term)
Working temperature	4°C to 30°C
Storage solution	20% ethanol
Storage temp	4°C to 8°C
Filter plate material	Polypropylene and polyethylene
Filter plate size <sup>3</sup>	127.8 × 85.5 × 30.6 mm
Volume, prepacked medium/well	50 μL
Wellvolume	800 µL
Centrifugation speed <sup>4</sup>	700 × g
Vacuum pressure <sup>4</sup>	
Recommended	-0.1 to -0.3 bar
Maximum	- 0.5 bar
1 =	

<sup>&</sup>lt;sup>1</sup> Protein dependent

## **Ordering information**

#### **Products**

Description	Quantity	Code No.
Protein A HP	4 × 96-well	28903133
MultiTrap	filter plates	

### **Related products**

Description	Quantity	Code No.
Sample Grinding Kit	50 samples	80648337

Description	Quantity	Code No.
Protease Inhibitor Mix	1 mL	80650123
Nuclease Mix	0.5 mL	80650142
NHS HP SpinTrap™	5 mL medium, 24 columns	28903128
Streptavidin HP SpinTrap	16 columns	28903130
Streptavidin HP	4×96-well	28903131
MultiTrap	filter plates	
Protein A HP SpinTrap	16 columns	28903132
Protein G HP SpinTrap	16 columns	28903134
Protein G HP	4×96-well	28903135
MultiTrap	filter plates	
Collection Plate	5 × 96 well plates	28403943
Ab SpinTrap	50 × 100 μL	28408347
Ab Buffer Kit	1	28903059
Protein A/G SpinTrap buffer kit	1	28913567

#### Literature

Code No.	
28906789	
18103746	
18102229	
	28906789 18103746

 $<sup>^2~{\</sup>rm pH}$  below 3 is sometimes required to elute strongly bound Ig species. However, protein ligands may hydrolyze at very low pH.

According to American Standard Institute (ANSI) and Society for Biomolecular Screening (SBS) standards 1-2004, 3-2004 and 4-2004.

 $<sup>^4\,</sup>$  Actual settings will depend on the sample properties and pretreatment.

## **Cue card: Antibody purification protocol**

## 1 Prepare collection plates

2 Remove storage solution		Shake gently upside down
		Remove the seals
		1 min
		700×g
B Equilibrate		Add 300 µL binding buffer
		30 s
		70-100 × g
4 Bind antibody	2004	Add 300 µL antibody in binding buffer
		Incubate 4 min on shaker
		30 s
		70-100 × g
5 Wash	2 ×	Add 300 µL binding buffer
		30 s
		70-100 × g
6 Elute antibody	2 ×	Add 200 µL binding buffer
	~~~~	30 s
		70 × g

## **Cue card: Cross-link protocol**

1 Remove storage solution		Shake gently upside down
		Remove the seals
		1 min
		700 × g
2 Equilibrate	3 ×	Add 400 µL binding buffer
		1 min
		700 × g
3 Bind antibody	262/	Add 200 µL antibody in binding buffer
		Incubate 30 min on shaker
		1 min
		700 × g
4 Wash	~~~~	Add 400 µL binding buffer
		1 min
		700 × g
5 Change buffer		Add 400 µL triethanolamine
		1 min
		700 × g
6 Cross-link		Add 400 µL DMP in triethanolamine
		Incubate 60 min on shaker
		1 min
		700 × g

7 Wash	~~~~/ ~~~/	Add 400 µL triethanolamine
		1 min
		700 × g
8 Block		Add 400 µL triethanolamine
	2001	Incubate 15 min
		1 min
		700 × g
9 Remove unbound antibody	2 ×	Add 400 µL elution buffer
		1 min
	44	700 × g
10 Wash	2 ×	Add 400 µL binding buffer
		1 min
		700 × g
11 Bind target protein	a≈07	Add 200 µL sample in binding buffer
		Incubate 60 min on shaker
		1 min
		700 × g
12 Wash	5 ×	Add 400 µL wash buffer
		1 min
		700 × g
13 Elute	3 ×	Add 200 µL elution buffer
		1 min
	797	1000 × g
	TO V	

## **Cue card: Classic protocol**

1 Remove storage solution		Shake gently upside down
		Remove the seals
		1 min
		700×g
2 Equilibrate	-001	Add 400 µL binding buffer
		1 min
	3 ×	700×g
3 Bind antibody	2621	Add 200 µL antibody in binding buffer
		Incubate 30 min on shaker
		1 min
		700 × g
4 Wash	ZÖ. X /	Add 400 µL binding buffer
		1 min
		700×g
Bind target protein	JK 07	Add 200 µL sample in binding buffer
		Incubate 60 min on shaker
		1 min
		700 × g

6 Wash	- TO	Add 400 μL wash buffer	
		1 min	
	5 ×	700 × g	
7 Elute		Add 200 μL elution buffer	
		Shake for 1 min	
		1 min	
	3×	700 × g	

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