

# **UptiSpin Protein A**

SECTION 1		SECTION 3	
Introduction	1	Easy-to-read protocols for purifying	
UptiSpin Protein A benefits	1	antibodies using Protein A	9
Contents of a UptiSpin Protein A column kit	2	Delipidation procedure	9
Additional recommended equipment	3	Buffers in the kit	9
Storage conditions	3	Step by step Mini protocol	10
Specifications	3	Step by step Midi protocol	11
SECTION 2		APPENDIX	
Principles of Protein A and G chromatography	4	Procedure to convert RPM to RCF in a swing	
Affinity chromatography	5	bucket rotor	13
General conditions for optimal binding	6	Procedure to convert RPM to RCF in a fixed	
Choosing the correct buffer conditions	6	angle rotor	13
Eluting the protein from an UptiSpin column	7	Protein A kit buffer formulation	14
Binding kinetics of UptiSpin A spin column	7	Questions and answers	14
Protein A affinity resin	7	Troubleshooting assistant	16
Choosing between Protein A and G	8	Glossary	17
ŭ		References	18
		Ordering information	18

Section 1

## Introduction:

Affinity purification of monoclonal antibodies has been largely confined to the use of Protein A and Protein G chromatography. The UptiSpin A kit is designed for simple, rapid antibody purification from serum, ascites and tissue culture supernatant such as those derived from static cultures and bioreactors. UptiSpin column replace lengthy and expensive chromatographic methods such as FPLC®

This UptiSpin antibody purification kit incorporates pre-packed Protein A resin plugs in ready-to-use spin column. The objective is to offer the researcher total protein purification solutions from the initial fractionation stage to the final polishing steps. Resolution of the monoclonal antibody is achieved either in a 2.2 ml centrifuge tube for the UptiSpin Mini column or in a 50 ml centrifuge tube for the UptiSpin Midi column. The rapid purification protocols provided in this handbook for affinity chromatography permit the recovery of high levels of pure antibodies in minutes. Large numbers of samples can be processed at the same time. There is negligible hold-up volume, ensuring high solute recovery with minimal non-specific absorptive losses. UptiSpin employs powerful resin-based technology for separating proteins and involves only a few steps, making the isolation of pure antibody samples rapid and simple to perform. Antibody samples purified using UptiSpin column may be used in a wide range of laboratory procedures such as 1D or 2D polyacrylamide gel electrophoresis, Western blotting, ELISA, immunohistochemical or immunofluorescence studies. The antibodies are sufficiently pure for radiolabelling, conjugations (for example fluorescein) or preparation of immunoaffinity column.

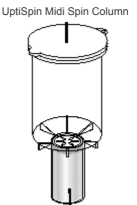
FPLC® is a registered trademark of Amersham Biosciences

# **UptiSpin Benefits:**

- UptiSpin A kits are designed to eliminate tedious chromatographic steps normally associated with Protein A chromatography.
- UptiSpin A reduces time-to-purity by incorporating protocols to suit various applications, supported by a comprehensive handbook and required buffers in a convenient kit format.
- The beaded supports offer excellent flow properties. This combined with the tapering of the spin column provides uniform flow paths that allow optimal use of the available resin bed in swing bucket rotors.
- Negligible hold-up volume of the Protein A resin plug ensures high antibody recovery.
- The provision of a disposable spin column is ideally suited to GMP production where current user requirements and price sensitivities demand regeneration of the affinity matrix using harsh treatments e.g. peroxyacetic acid for Protein A or G resins.





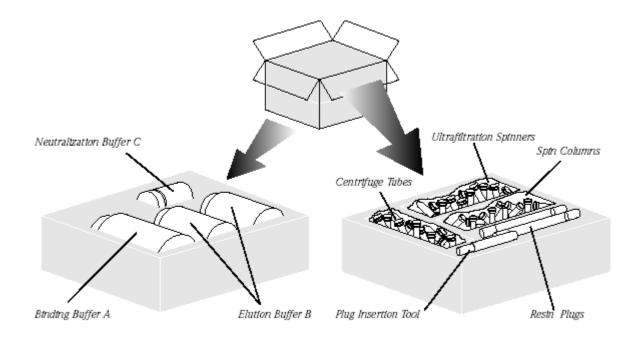


UptiSpin Protein A Midi Plug

# UptiSpin Protein A Mini spin column kit contents:

The UptiSpin Mini kit contains:

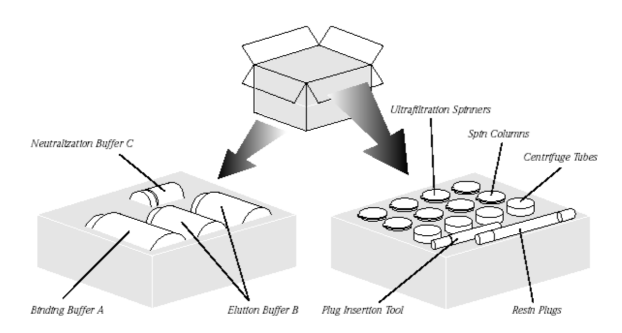
- 16 x UptiSpin column plugs containing Protein A-agarose resin.
- 16 x UptiSpin column barrels (0.65 ml capacity in a fixed angle rotor).
- 32 x 2.2 ml microcentrifuge tubes.
- 16 x 10 kDa MWCO (Regenerated cellulose) ultrafiltration spinners.
- 1 x 250 ml binding buffer (buffer A) bottle.
- 1 x 125 ml elution buffer (buffer B1) bottle.
- 1 x 125 ml elution buffer (buffer B2) bottle.
- 1 x 30 ml neutralization buffer (buffer C) bottle.
- Plug insertion tool.
- · Comprehensive handbook.
- · Ultrafiltration spinner instruction sheet.
- · Protocol card.



# UptiSpin Protein A Midi spin column kit contents :

The UptiSpin Midi kit contains:

- 4 x UptiSpin column plugs containing Protein A-agarose resin.
- 4 x UptiSpin column devices (20 ml capacity in a swing bucket rotor).
- 8 x 50 ml centrifuge tubes.
- 4 x 30 kDa MWCO 15 ml (Regenerated cellulose) ultrafiltration spinners.
- 1 x 250 ml binding buffer (buffer A) bottle.
- 1 x 125 ml elution buffer (buffer B1) bottle.
- 1 x 125 ml elution buffer (buffer B2) bottle.
- 1 x 30 ml neutralization buffer (buffer C) bottle.
- · Plug insertion tool.
- · Comprehensive handbook.
- Ultrafiltration spinner instruction sheet.
- Protocol card.





## Additional equipment recommended:

- Filters units : 0.2 and 1.2  $\mu$ m syringe filters for clarification. Our recommended filter unit is the 0.2 mm Steriflip GP unit from Millipore Corp (Cat. No : SCGP 005 25).
- Quartz cuvettes for UV absorbance measurements.
- UV/VIS spectrophotometer.
- Pasteur pipettes and micro-pipettes.
- · Marker pen and test tube rack.
- A bench-top centrifuge with swing bucket rotor that can accommodate 50 ml centrifuge tubes.
- A microcentrifuge that can accommodate 2.2 ml microcentrifuge tubes.

N.B. (For the Midi spin column only). The preferred rotor is a **swing bucket** rotor. For optimal performance with a fixed angle rotor, ensure that the orientation of the spin column in the rotor is the same for sample binding, washing and elution steps.

## Storage conditions:

Remove the UptiSpin Protein A plug box from the kit and store it at 2-8°C. There is no need to place the rest of the kit in a fridge or cold room. All buffers, for example, contain 0.1% sodium azide and can be stored at room temperature. Do not freeze the resin plugs. Freezing the suspension will damage the agarose beads. The resin is pre-swollen and de-fined. It is transported in 0.01% Thimerosal. UptiSpin column are stable for up to 2 years at 2-8°C from the date of manufacture. Expiry date is recorded clearly on the outside of the pack.

## Disclaimer:

This product is for research use only and is not intended for use in clinical diagnosis. No claims beyond replacement of unacceptable material or refund of purchase price shall be allowed.

### **Specifications:**

Source Recombinant Protein A expressed in E. coli (N.B. No toxic bacterial contaminants

normally found in native Protein A)

Supporting UptiSpin matrix Covalently coupled to agarose resin

Max sample volume 20 ml (Midi, Swing bucket rotor), 0.65 ml (Mini, Fixed angle rotor)

Resin bed volume1.6 ml (Midi) 0.23 ml (Mini)Ligand density3,5 mg Protein A/ml resin

**Bead size range** 60-165 μm **Recommended working pH** pH2.5-9.0

Typical binding capacity ≥ 20 mg rabbit lgG (Midi)

≥ 1 mg rabbit lgG (Mini)

(Note that the Protein A resin can have different binding capacities to subtypes of

immunoglobulins derived from the same species).

Leaching levels < 5 ng recombinant Protein A/ml

Chemical stability High
Endotoxin levels Unknown

**Toxin levels** Free of Staphyloccocus enterotoxins and hemolysins

Solubility in water Insoluble
Colour coded end-caps Red

Section 2

# Chemical compatibility of the UptiSpin column :

All resins are susceptible to oxidative agents. Avoid high temperatures. The spin columns have high chemical resistance to short exposure to organic solvents (e.g. 70% ethanol, 5.8 M acetic acid) and are stable in all aqueous buffers commonly used for Protein A chromatography. Protein A is resistant to 6 M guanidine-HCI, 8 M urea and 2 M sodium isothyocyanate.

# Immuno-affinity separations:

Three types of immunoaffinity ligands are commonly employed:

- (i) Protein A & G.
- (ii) Antibody directed against the monoclonal species (usually anti-mouse IgG).
- (iii) Antigen to which the monoclonal antibody was raised.



## Principles of Protein A and G chromatography:

This handbook focuses specifically on Protein A affinity chromatography. All modes of chromatography can be used effectively for the separation of antibodies. Although ion exchange chromatography can resolve different polyclonal antibodies and different subclasses, a degree of customization of the protocol is required. Affinity techniques include Protein A or G, immobilized anti-antibodies and immobilized antigens. The simplicity of Protein A is extremely attractive as it lends itself to the bind, wash and elute mode of operation if the appropriate buffer formulations are selected. The use of Protein A and G is widespread and has largely superseded the use of anti-antibodies. Protein A chromatography therefore, remains the principal theme of this handbook.

## Application drivers for Protein A and G chromatography:

Monoclonal antibody purification for solution state assays and Western blotting	Mini & Midi
Semi-preparative antibody purification	Midi
Purification of polyclonal antibodies for solution state assays and Western blotting	Mini & Midi
Removal of endotoxins from an antibody solution	Mini & Midi
Small scale antibody purification	Mini
Screening antibody expression	Mini
Immunoprecipitation studies	Mini
Purification of antibodies for immuno-histochemical and immunofluorescence studies	Mini

### Affinity chromatography:

The essence of affinity chromatography utilizes the concept of bio-specificity, implying an interaction between a natural binding site and the natural ligand, whether it be enzyme-substrate, enzyme-inhibitor or Protein A/G-antibody interactions. Often a researcher needs to choose the correct base matrix, select the optimal activation chemistry and couple a suitable ligand to develop an affinity column to purify the target protein. UptiSpin column remove the guesswork associated with optimizing the resin chemistry by offering ready-to-use resin plugs that have satisfied stringent quality control to guarantee reproducible purification performance. Many affinity support preparations require less than 5 ml resin so these Mini and Midi spin column contain ideal bed volumes.

The only decisions required by the researcher are whether they wish to work with Protein A or G resin. Optimal buffer and elution conditions for the purification step of many common IgG subclasses have been defined and these are supplied with the kits.

Most polymeric mini-column currently offered operate by gravity flow and typically contain less than 5 ml resin. The column are normally constructed from polystyrene or polypropylene with a simple barrel and the addition of porous disks to contain the resin. However, the affinity separation often takes several hours to complete; in addition the researcher usually has to pack the column, which can add a minimum 2 further hours to the purification step. The elaborate nature of the packing means that it is usually undertaken by an experienced laboratory worker.

The majority of purifications require fast processing times to minimise the hydrolytic actions of proteases. UptiSpin Protein A kits allow multiple parallel purifications to be achieved without the need to employ expensive PEEK tubing (protein-friendly)- based chromatography systems. For example, 12 parallel small scale antibody purifications can be achieved in a microfuge containing a 12 tube fixed angle rotor.

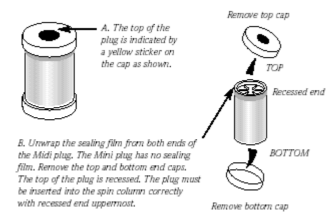


Fig. 1. Schematic diagram of a UptiSpin Mini or Midi spin column plug.

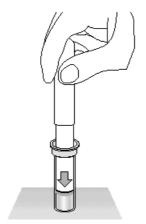
The plug and play concept for the UptiSpin Mini spin column



Top of plug

Loading the plug into the spin

Place the plug into the spin column with the recessed end uppermost.

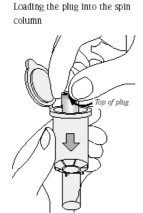


Push the plug FULLY into the tapered end of the spin column using the plug insertion tool.



It is now ready for pre-equilibration with binding buffer followed by centrifugation.

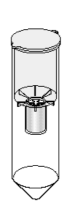
# The plug and play concept for the UptiSpin Midi spin column



Place the plug into the spin column with the recessed end uppermost.



Push the plug FULLY into the tapered end of the spin column using the plug insertion tool.



It is now ready for preequilibration with binding buffer followed by centrifugation.



After use, the plug is removed using the plug insertion tool.

## General considerations for selecting optimal binding conditions for UptiSpin Protein A spin column:

Any sample, such as a crude biological extract, a cell culture supernatant, serum, ascites or an artificial standard can be used in the UptiSpin A spin column. It is important that the sample is first filtered through a **0.2 mm** filter to remove particulates that could clog the resin flow channels. All samples should be filtered just prior to loading even if they have been filtered several days before the chromatographic run. Aggregation/precipitation of proteins is common during storage and repeated freeze/thaw cycles in sera, ascites and tissue culture supernatants. Lipids, which can be found at high levels in serum or ascites should also be removed (see delipidation protocol on page 9). Of equal importance is the ability to process the samples rapidly and, if the need arises, to be able to purify the target protein in a UptiSpin column at 4°C.

Optimal conditions for binding the target molecule to a resin are critical for successful separation of the protein. If the binding conditions are not optimal with respect to pH, salt concentration, presence or absence of metal ions etc, the interaction could be weak or non-existent. In many instances, the sample needs to be dialysed or diafiltered by ultrafiltration before it is applied to an affinity or ion exchange chromatographic support. In Protein A or G separations, the sample should simply be diluted 1:1 (v/v) in  $1 \times 1 \text{ x}$  binding buffer provided in the kit.

The interaction of immobilized Protein A or Protein G with immunoglobulins (Igs) is pH-dependent. The binding capacity for Protein A is optimal at pH 8-9, whereas the binding capacity of Protein A is high over a broader pH range. Salt concentration can significantly affect the binding of mouse Igs to Protein A by reducing severe ionic interactions and enhancing hydrophobic interactions. Mouse IgG1, rat IgG1 and rat IgG2b bind well to immobilized Protein A when the salt concentration is higher than 1 M, but bind poorly at low salt concentrations. Note, however, that their binding capacities can be substantially lower than those antibodies that bind strongly to Protein A or G resin.

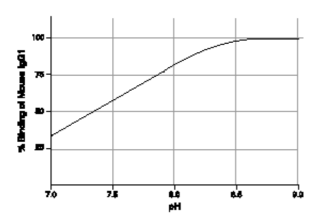


Fig. 2. Typical pH dependence of binding of mouse IgG1 to Protein A

## Choosing the correct buffer conditions for Protein A separations:

Typical binding buffers employed in Protein A affinity separations are :

- (i) 1-1.5 M Glycine/NaOH, 2-3 M NaCl pH 9.0
- (ii) 1 M Sodium Borate, 2 M NaCl pH 9.0
- (iii) 0.1 M Sodium phosphate, 0.1 M NaCl pH 7.4 (PBS)

Typical elution buffers employed in Protein A affinity separations are :

- (i) 0.1 M Sodium citrate pH 3.0-6.0
- (ii) 0.1-0.2 M Glycine/HCl pH 2.5-3.0
- (iii) 0.1 M Sodium phosphate pH 3.0-6.0

The typical neutralization buffer used for Protein A affinity separations is :

(i)1M Tris/HCI pH9.0

# Eluting the antibody from a UptiSpin A column :

The most common elution conditions for Protein A or G affinity and immuno-affinity separations involve a reduction in pH to pH 2.5 (see *table 1*). It is important to appreciate that a few proteins (e.g. some monoclonal antibodies) are acid-labile and they can lose their activity at very low pH values. Above all, the elution conditions must preserve the integrity and activity of the target protein. Most observed denaturation is caused by harsh elution conditions. Acidic pH is known to reduce the antibody titre, decrease immuno-reactivity and distort the antibody structure. It is therefore critical that the pH is restored to neutrality after elution.

Subclass	Binding pH	Elution pH
IgG1	8.5-9.0	6.0-7.0
lgG2a	8.0-9.0	4.5-5.5
lgG2b	8.0-9.0	3.5-4.5
IgG3	8.0-9.0	4.0-7.5
IgG1	8.0-9.0	6.0-8.0
lgG2a	9.0	7.5-9.0
lgG2b	8.0-9.0	7.0-8.0
IgG2c	8.0-9.0	3.0-7.0
IgG1	7.0-7.5*	2.5-4.5
lgG2	7.0-7.5*	2.5-4.5
IgG3	7.0-7.5*	3.0-7.0
lgG4	7.0-7.5*	2.5-4.5
IgG	7.5	3.0-7.0
IgG1	7.5-9.0	4.0-5.0
IgG2	7.5-8.0	3.0-4.5
	IgG1 IgG2a IgG2b IgG3 IgG1 IgG2a IgG2b IgG2c IgG2c IgG1 IgG2 IgG3 IgG4 IgG	IgG1     8.5-9.0       IgG2a     8.0-9.0       IgG2b     8.0-9.0       IgG3     8.0-9.0       IgG1     8.0-9.0       IgG2a     9.0       IgG2b     8.0-9.0       IgG2c     8.0-9.0       IgG1     7.0-7.5*       IgG2     7.0-7.5*       IgG3     7.0-7.5*       IgG4     7.5-7.5*       IgG     7.5       IgG1     7.5-9.0

## Binding kinetics of UptiSpin A spin column:

The flow rate through an affinity chromatography support is important in achieving optimal separation. Flow rate through the column support is inextricably related to the efficiency of the separation; too fast a flow will cause the mobile phase to move past the beads faster than the diffusion time necessary to reach the internal bead volume. Our studies demonstrate that the large internal surface area of the UptiSpin Midi resin bed compensates completely for the velocity of the mobile phase through the column support when the centrifugal speed does not exceed 1,500 g\*.

\* No UptiSpin Midi Protein A and G performance data is available for centrifugal speeds greater than 1,500 g. Centrifugal speeds as high as 13,000 rpm (equivalent to 11,960 in a fixed angle rotor with an average radius of 49 mm) have been achieved with he UptiSpin Protein A and G Mini spin column. The Protein A resin chemistries used in the UptiSpin column have sufficiently rapid association kinetics between the protein molecule and the immobilized ligand to

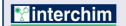


Table 1.

allow for optimal diffusional flow through the internal bead structure. Traditionally, gravity flow chromatography is very slow and resolution of the protein separation can be adversely affected by secondary diffusion effects.

Source	Туре	Total Ab	Specific Ab	Contamination
Serum	Polyclonal	10 mg/ml	1 mg/ml	Other serum Ab
Stattc Culture	Monoclonal	1 mg/ml	0.05 mg/ml	Medium serum Ab
Bioreactor	Monoclonal	10 mg/ml	9 mg/ml	Medium serum Ab
Ascites	Monoclonal	10 mg/ml	9 mg/ml	Mouse Ab

## Protein A affinity resin:

Protein A is a cell wall protein from *Staphylococcus aureus* with a molecular weight between 35-50 kDa. The quality of the Protein A agarose is important to avoid leakage of Protein A during the elution procedure. Immobilized Protein A resins linked via an amide bond to the UptiSpin between the amino groups of Protein A or G and either oxirane or N-hydroxysuccinimide ester groups form the most stable cross-links. Immobilized Protein A binds specifically to the Fc region of immunoglobulin molecules of many mammalian species.

Protein A affinity chromatography is a rapid one-step purification, which removes most non-IgG contaminants and can achieve purities close to homogeneity. It is particularly useful for purifications of tissue culture supernatant, where 10-100 fold concentrations can be achieved.

Key code for relative affinity of Protein A & G for respective antibodies:
 ✓✓✓ = Strong affinity
 ✓✓ = Moderate affinity
 ✓✓ = Weak affinity
 ✓ = Slight affinity
 X = No affinity

# Binding Affinities of Protein A and Protein G

	Protein A	Protein G		Protein A	Protein G
Human IgG1	<b>VVVV</b>	<b>VVVV</b>	Rabbit IgG	VVVV	VVV
Human IgG2	VVVV	<b>VVVV</b>	Hamster IgG	~	VV
Human IgG3	×	<b>VVVV</b>	Guinea Pig IgG	<b>VVVV</b>	VV
Human IgG4	VVVV	<b>VVVV</b>	Bovine IgG	VV	VVVV
Human IgA	VV	×	Sheep IgG	<b>√</b> / <b>×</b>	VV
Human IgD	VV	×	Goat IgG	<b>√</b> / <b>×</b>	VV
Human IgE	VV	×	Ptg IgG	VVV	VVV
Human IgM	VV	×	Chicken IgG	×	~
Mouse IgG1	~	VV			
Mouse IgG2a	VVVV	<b>VVVV</b>	Fragments		
Mouse IgG2b	VVV	VVV			
Mouse IgG3	~~	VVV	Human Fab	~	~
Mouse IgM	<b>√</b> / <b>×</b>	×	Human F(ab')2	~	~
Rat IgG1	*	~	Human scFv	~	*
Rat IgG2a	×	<b>VVVV</b>	Human Fc	VV	VV
Rat IgG2b	*	~	Human ĸ	×	*
Rat IgG2c	~	VV	Human $\lambda$	×	×
Rat IgM	<b>√</b> / <b>×</b>	×			

## Choosing between Protein A and Protein G UptiSpin columns

Immunoglobulin G from most species consists of several subclasses with different biological properties. Four subclasses of IgG have been identified in human (IgG1, IgG2, IgG3, and IgG4) and in mouse (IgG1, IgG2a, IgG2b and IgG3). For immunological studies, it is often necessary to isolate one particular subclass of IgG from the other subclasses. Protein G binds to all major Ig classes except IgM and therefore has a wider reactivity profile than Protein A. However, the binding of Igs to Protein A is often stronger, making elution and complete recovery of the immunoglobulin more difficult. Interestingly, due to the lower cost of Protein A compared to Protein A, researchers tend to experiment first with Protein A followed by Protein G. Protein A withstands harsher conditions used in cleaning-in-place procedures. However, UptiSpin column obviate the need for cleaning.

The affinity of interaction of Protein A with mouse IgG subclasses varies. The most common subclass of mouse monoclonal antibodies is IgG1. Customization of the purification strategy may be required for the affinity separation as mouse IgG1 does not generally bind well to protein A. However, as the affinity interaction is pH- and salt dependent, under high salt regimes (2-3 M NaCl) and high pH (pH 8-9), the antibodies will bind to Protein A.

The needs of the researcher dictate that the speed of sample processing, the cost and the reproducibility are key criteria for selecting purification tools. By selecting immunoglobulin-binding proteins with the appropriate cross-reactivities coupled to careful design of the strategic protocol, objectives such as the selective recovery of therapeutic antibodies from complex mixtures is assured.

lmmunoglobulin Heavy chain	IgG1 71	lgG2 %	IgG3 ≇₃	lgG4 74	$_{\mu}^{\mathrm{IgM}}$	IgA1 γι	$^{\rm lgA2}_{\alpha_1}$	stgA $\alpha_1$ or $\alpha_2$	IgD δ <sub>l</sub>	lgE ε <sub>1</sub>
Mean serum	9	3	1	0.5	1.5	3.0	0.5	0.05	0.03	0.0000
concentration (mg/ml)										
Sedimentation	7S	7S	75	7S	19S	7S	75	11S	7S	88
constant	, 5	13	10	10	195	13	10	113	13	0.0

Table 2. Physical properties of immunoglobulin subclasses.

lmmunoglobulin	IgG1	lgG2	IgG3	lgG4	IgM	IgA1	lgA2	sigA	IgD	IgE	
Heavy chain	$\gamma_1$	1/2	$\gamma_3$	$\gamma_4$	$\mu$	$\gamma_1$	$\alpha_1$	$\alpha_1$ or $\alpha_2$	$\delta_1$	$\epsilon_1$	
Mean serum	9	3	1	0.5	1.5	3.0	0.5	0.05	0.03	0.00005	
concentration (mg/ml)											
Sedimentation	7S	7S	7S	7S	19S	7S	7S	11S	7S	85	
constant	15 15		1.0	,,,	1,0		10			00	
Molecular weight	146,000	146,000	170,000	146,000	970,000	160,000	160,000	385,000	184,000	188,000	
Molecular weight	51.000	£1,000	60.000	£1,000	65,000	F4 000	F2 000	53.54.000	60 700	70.500	
of heavy chain	51,000	51,000	60,000	51,000	65,000	56,000	52,000	52-56,000	69,700	72,500	
Number of heavy	4	4	4	4	5	4	4	4	4	5	
chain domains	_	_		7	-	-		7	_	_	
% Carbohydrate	2-3	2-3	2-3	2-3	12	7-11	7-11	7-11	9-14	12	

Section 3

# Protocol for purifying monoclonal antibodies using UptiSpin Protein A kits

Immunoglobulin binding proteins have been used extensively for the purification of IgG molecules from serum, ascites, or cell culture supernatants such as those derived from static cultures and bioreactors. The affinity of Protein A for the Fc region of immunoglobulins makes it the natural ligand choice for many researchers involved in IgG isolation. We recommend the following elution buffers in Tables 3 (3A (Mini) and 3B (Midi)) with the respective subclass of antibodies.

## **Delipidation procedure:**

All protein A & G affinity column are affected by the presence of lipids and lipoproteins, especially in antibody samples derived from ascites fluid. For end users who have antibody solutions which they need to delipidate, the following protocol is a gentle and easy method for removing lipids and lipoproteins.

- 1) Add 0.04 ml 10% dextran sulphate solution and 1 ml 1 M calcium chloride per ml sample.
- 2) Mix for 15 min.
- 3) Centrifuge at 10,000 g for 10 min.
- 4) Discard the precipitate.
- 5) Exchange the sample into TBS (Tris Buffered Saline) using dialysis, ultrafiltration or a desalting column. Do not buffer exchange into a phosphate-containing buffer such as PBS.

## Buffers in the kit:

All buffers contain 0.1% sodium azide as a preservative.

- Binding buffer (Buffer A):
- 0.1 M sodium phosphate buffer, 0.15 M NaCl, pH 7.4
- Elution buffer (Buffer B2) :
- 0.2 M Glycine/HCl buffer pH 2.5
- Neutralization buffer (Buffer C):
- 1 M Tris/HCl buffer pH 9.0

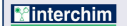


Table 3A. Mini spin columns :

Choice of Binding, Elution and Neutralization Buffers for the following antibody subclasses and host species.

Species	Subclass	Binding buffer	Elution buffer	Neutralization buffer (NB)
Mouse	lgG1	Buffer A	Buffer B1 ———	Buffer C (add 25 $\mu\text{l}$ NB to 0.5 ml final eluate) followed by B2
	.5		Buffer B2	Buffer C (add 65 μl I NB to 0,5 ml final eluate)
Mouse	lgG2a	Buffer A	Buffer B2	Buffer C (add 65 μl NB to 0,5 ml final eluate)
Mouse	lgG2b	Buffer A	Buffer B2 ———	Buffer C (add 65 μl NB to 0,5ml final eluate)
Mouse	lgG3	Buffer A	Buffer B2	Buffer C (add 65 μl NB to 0,5 ml final eluate)
			Buffer B1 ———	Buffer C (add 25 $\mu$ l NB to 0,5 ml final eluate) followed by B2
Rat	lgG1	Buffer A		
			Buffer B2 ———	Buffer C (add 65 μl NB to 0,5 ml final eluate)
			Buffer B1 ———	Buffer C (add 25 μl NB to 0,5 ml final eluate) followed by B2
Rat	IgG2a	Buffer A	D " D0 -	
			Buffer B2 ——	Buffer C (add 65 μl NB to 0,5 ml final eluate)
			Buffer B1 ——	Buffer C (add 25 µl NB to 0,5 ml final eluate) followed by B2
Rat	lgG2b	Buffer A	bullel b I — =	Bullet C (add 25 $\mu$ i NB to 0,5 iii iiilal eldate) lollowed by B2
rtat	19025	Ballet /	Buffer B2 ——	Buffer C (add 65 μl NB to 0,5 ml final eluate)
Rat	lgG2c	Buffer A	Buffer B2	Buffer C (add 65 µl NB to 0,5 ml final eluate)
Human	lgG1	Buffer A	Buffer B2 ———	Buffer C (add 65 µl NB to 0,5 ml final eluate)
Human	lgG2	Buffer A	Buffer B2	Buffer C (add 65 µl NB to 0,5 ml final eluate)
Human	IgG3	Buffer A	Buffer B2 ———	Buffer C (add 65 µl NB to 0,5 ml final eluate)
Human	IgG4	Buffer A	Buffer B2 ———	Buffer C (add 65 µl NB to 0,5 ml final eluate)
Rabbit	IgG	Buffer A	Buffer B2	Buffer C (add 65 µl NB to 0,5 ml final eluate)
Guinea pig	IgG1	Buffer A	Buffer B2	Buffer C (add 65 µl NB to 0,5 ml final eluate)
Guinea pig	lgG2	Buffer A	Buffer B2 ———	Buffer C (add 65 µl NB to 0,5 ml final eluate)
1 0	•			, , , , , , , , , , , , , , , , , , , ,

For all other Igs e.g. Hamster IgG, use elution buffer B2 only.

## Step by step protocol for Mini UptiSpin Column

#### **RESIN PLUG LOADING**

1. Load the pre-packed resin Mini plug containing immobilized recombinant Protein A resin into the barrel of the UptiSpin column using the insertion tool as shown on page 5.

#### PRE-EQUILIBRATION (Total spin time = 2 min)

- 2. Equilibrate the Protein A spin column with 0.65 ml binding buffer A, pH 9 by centrifuging the spin column at 1,800 g (4,400 rpm in a Heraeus Biofuge Pico or 5,000 rpm in a Sanyo MSE Micro Centaur) for 1 min\*. Repeat this preequilibration step with 0.65 ml binding buffer A, pH 9 at 1,800 g for 1 min.
- \* If 1 spin column is to be used, ensure that the spin column is counterbalanced in the microfuge with a microcentrifuge tube filled with the correct level of water.

## CLARIFICATION OF SAMPLE

- 3. Filter 1 ml sample through a single 0.2 µm syringe filter to remove any cellular debris.
- N.B : Protein precipitation is common during storage and repeated freeze/thaw cycles in ascites, sera and tissue culture supernatants. As with all forms of chromatography, it is important that the sample is filtered through a final 0.2 μm syringe filter **immediately** before loading it on to the spin column.

# SAMPLE LOADING (Total spin time = 6 min)

4. Dilute the sample 1:1 (v/v; eg. add 0.5ml 0.2 µm filtered sample to 0.5ml binding buffer A, pH 9). Mix by inverting the capped tube 3-4 times. Pipette the 0.65ml sample into the spin column. Centrifuge the spin column at 640g (2,600 rpm in a Heraeus Biofuge Pico or 3,000 rpm in a Sanyo MSE Micro Centaur) for 6 min.

N.B: Increase the spin time or speed if any sample remains above the plug.

## WASHING (Total spin time = 3 min)

5. Wash the spin column three times with 0.65ml binding buffer A, pH 9 to remove unbound contaminants by centrifuging the UptiSpin column for 1 mn at 1,800g (4,400 rpm in a Heraeus Biofuge Pico or 5,000 rpm in a Sanyo MSE Micro Centaur). The unbound wash will contain non-immunoglobulin components.



ELUTION (Total spin time = 4 min)
For purifying mouse IgG1, rat IgG1, rat IgG2a, rat IgG2b and bovine IgG1, use both elution steps 6 and 7.

ELUTION (Total spin time = 4 min)
For purifying unassigned IgG, mouse IgG2a, mouse IgG2b, mouse IgG3, rat IgG2c, human IgG1-IgG4, rabbit IgG, guinea pig IgG1, guinea pig IgG2, bovine IgG2 and any other IgGs, proceed to elution step 7 only.

6. Elute the bound IgG with 0.5ml elution buffer B1 directly into a fresh centrifuge tube containing 25µl neutralization buffer C to bring the pH of the sample to approximately 7.5. Centrifuge the UptiSpin column for 2min at 1,800g. Swirl the tube to ensure thorough mixing of the final eluate with neutralization buffer C. Repeat this elution step.

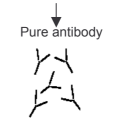
N.B: Do not pool the two eluate fractions if you want to recover *concentrated* purified antibody.

7. Elute the bound IgG with 0.5ml elution buffer B2 directly into a fresh centrifuge tube containing 65 µl neutralization buffer C to bring the pH of the sample to approximately 7.5. Centrifuge the UptiSpin column for 2min at 1,800g. Swirl the tube to ensure thorough mixing of the final eluate with neutralization buffer C. Repeat this elution step.

N.B: Do not pool the two eluate fractions if you want to recover *concentrated* purified antibody.

# DESALTING AND CONCENTRATING THE PURIFIED ANTIBODY

8. If necessary, de-salt and concentrate the antibody preparation using the 10kDa MWCO ultrafiltration spinner supplied. Add 0.05-0.2% w/v sodium azide if the antibodies are to be stored at 2-8°C. We recommend freezing the antibodies in small aliquots in 10-50% glycerol at -20°C for long term storage.



REGENERATION OF THE PROTEIN A MINI PLUG

9. Wash the Mini plugs twice with 0.65ml elution buffer B2 (pH 2.5) by centrifuging the spin column at 1,800g for 1min. Then wash the plugs twice with 0.65ml binding buffer A (pH 9.0) by centrifuging the spin column at 1,800 g for 2min. Proceed to the pre-equilibration step of another bind-wash-elute cycle if the plugs are to be re-used immediately. After regeneration, plugs can also be stored, without their end caps, in binding buffer A or in 0.1% sodium azide (made up in distilled water) at 2-8°C until further use.





## Easy-to-read Mini Purification Protocol E.g. Human serum

Fraction	Volume	Step	RCF	Spin Time
Pre-equilibration #1	0.65 ml	BBA pH 9.0	1,800 g	1 min
Pre-equilibration #2	0.65 ml	BBA pH 9.0	1,800 g	1 min
Sample Loading	0.65 ml	1:1 serum: BBA pH 9.0	640 g	6 min
Wash #1	0.65 ml	BBA pH 9.0	1,800 g	2 min
Wash #2	0.65 ml	BBA pH 9.0	1,800 g	2 min
Wash #3	0.65 ml	BBA pH 9.0	1,800 g	2 min
Final Eluate #1	0.5 ml	EB2 65 μl NBC	1,800 g	2 min
Final Eluate #2	0.5 ml	EB2 - 65 μl NBC	1,800 g	2 min

# **Easy-to-read Mini regeneration Protocol**

Fraction	Volume	Step	RCF	Spin Time
Clean-up #1	0.65 ml	EB2 pH 2.5	1,800 g	2 min
Clean-up #2	0.65 ml	EB2 pH 2.5	1,800 g	2 min
Wash #1	0.65 ml	BBA pH 9.0	1,800 g	2 min
Wash #2	0.65 ml	BBA pH 9.0	1,800 g	2 min

Table 3B. Midi spin columns:

Choice of Binding, Elution and Neutralization Buffers for the following antibody subclasses and host species.



Species	Subclass	Binding buffer	Elution buffer	Neutralization buffer (NB)
			Buffer B1 ———	Buffer C (add 0.5 ml NB to 10 ml final eluate) followed by B2
Mouse	lgG1	Buffer A		
		D " A	Buffer B2	Buffer C (add 1.3 ml NB to 10 ml final eluate)
Mouse	lgG2a	Buffer A	Buffer B2 ———	Buffer C (add 1.3 ml NB to 10 ml final eluate)
Mouse	lgG2b	Buffer A	Buffer B2 ———	Buffer C (add 1.3 ml NB to 10ml final eluate)
Mouse	lgG3	Buffer A	Buffer B2 ———	Buffer C (add 1.3 ml NB to 10 ml final eluate)
			Buffer B1 ———	Buffer C (add 0.5 ml NB to 10 ml final eluate) followed by B2
Rat	IgG1	Buffer A		
			Buffer B2 ———	Buffer C (add 1.3 ml NB to 10 ml final eluate)
			Buffer B1	Buffer C (add 0.5 ml NB to 10 ml final eluate) followed by B2
Rat	lgG2a	Buffer A		
	•		Buffer B2 ——	Buffer C (add 1.3 ml NB to 10 ml final eluate)
			Buffer B1 ——	Buffer C (add 0.5 ml NB to 10 ml final eluate) followed by B2
Rat	lgG2b	Buffer A	24.10. 2.	
1.00	19025	Danot 70	Buffer B2	Buffer C (add 1.3 ml NB to 10 ml final eluate)
Rat	lgG2c	Buffer A	Buffer B2 ———	Buffer C (add 1.3 ml NB to 10 ml final eluate)
Human	IgG1	Buffer A	Buffer B2	Buffer C (add 1.3 ml NB to 10 ml final eluate)
Human	lgG2	Buffer A	Buffer B2	Buffer C (add 1.3 ml NB to 10 ml final eluate)
Human	lgG3	Buffer A	Buffer B2	Buffer C (add 1.3 ml NB to 10 ml final eluate)
Human	lgG4	Buffer A	Buffer B2	Buffer C (add 1.3 ml NB to 10 ml final eluate)
Rabbit	IgG	Buffer A	Buffer B2	Buffer C (add 1.3 ml NB to 10 ml final eluate)
Guinea pig	lgG1	Buffer A	Buffer B2 ——	Buffer C (add 1.3 ml NB to 10 ml final eluate)
Guinea pig	lgG2	Buffer A	Buffer B2	Buffer C (add 1.3 ml NB to 10 ml final eluate)
Cantoa pig	.902	Danoi / C	Danoi DZ =	Editor & (add 1.6 fill 112 to 16 fill fillar bladte)

For all other Igs e.g. Hamster IgG, use elution buffer B2 only.

## Step by step protocol for Midi UptiSpin Column

#### **RESIN PLUG LOADING**

1. Load the pre-packed resin Midi plug containing immobilized recombinant Protein A resin into the barrel of the UptiSpin column using the insertion tool as shown on page 6.

## PRE-EQUILIBRATION (Total spin time = 3 min)

- 2. Equilibrate the Protein A spin column with 10ml binding buffer A, pH9.0 by centrifuging the spin column at 500g for 3min\*.
- \* If 1 spin column is to be used, ensure that the spin column is counterbalanced with a unit of equal weight (adjusted with distilled water) and without a Protein A resin plug.

# CLARIFICATION OF SAMPLE

- 3. Filter 12-15 ml sample through a single  $1.2\mu m$  (25mm diameter) syringe filter to remove any cellular debris. Then, filter the partially clarified sample through a single  $0.2\mu m$  (25mm diameter) syringe filter.
- N.B: Protein precipitation is common during storage and repeated freeze/thaw cycles in ascites, sera and tissue culture supernatants. As with all forms of chromatography, it is important that the sample is filtered through a final 0.2 µm syringe filter **immediately** before loading it onto the spin column.

## SAMPLE LOADING (Total spin time = 30min)

4. Dilute the sample 1:1 (v/v; eg. add 10ml 0.2µm filtered sample to 10ml binding buffer A, pH 9.0). Mix by inverting the capped tube 3-4 times. Pipette the 20ml sample into the spin column. Centrifuge the spin column at 100g for 30min.

N.B: Increase the spin time or speed if any sample remains above the plug.

## WASHING (Total spin time = 6min)

5. Wash the spin column twice with 10ml binding buffer A, pH 9.0 to remove unbound contaminants by centrifuging the UptiSpin column for 3min at 500g. The unbound wash will contain non-immunoglobulin components.

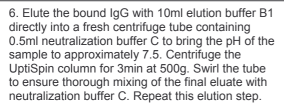
## ELUTION (Total spin time = 6min)

For purifying mouse IgG1, rat IgG1, rat IgG2a, rat IgG2b and bovine IgG1, use both elution steps 6 and 7.

ELUTION (Total spin time = 6min)

For purifying unassigned IgG, mouse IgG2a, mouse IgG2b, mouse IgG3, rat IgG2c, human IgG1-IgG4, rabbit IgG, guinea pig IgG1, guinea pig IgG2, bovine IgG2 and any other IgGs, proceed to elution step 7 only.





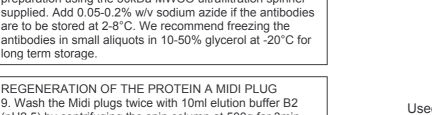
N.B: Do not pool the two eluate fractions if you want to recover concentrated purified antibody.

7. Elute the bound IgG with 10ml elution buffer B2 directly into a fresh centrifuge tube containing 1.3ml neutralization buffer C to bring the pH of the sample to approximately 7.5. Centrifuge the UptiSpin column for 3min at 500g. Swirl the tube to ensure thorough mixing of the final eluate with neutralization buffer C. Repeat this elution step.

N.B: Do not pool the two eluate fractions if you want to recover concentrated purified antibody.

## DESALTING AND CONCENTRATING THE PURIFIED **ANTIBODY**

8. If necessary, de-salt and concentrate the antibody preparation using the 30kDa MWCO ultrafiltration spinner supplied. Add 0.05-0.2% w/v sodium azide if the antibodies are to be stored at 2-8°C. We recommend freezing the antibodies in small aliquots in 10-50% glycerol at -20°C for long term storage.



9. Wash the Midi plugs twice with 10ml elution buffer B2 (pH2.5) by centrifuging the spin column at 500g for 3min. Then wash the plugs twice with 10ml binding buffer A (pH9.0) by centrifuging the spin column at 500 g for 3min. Proceed to the pre-equilibration step of another bind-washelute cycle if the plugs are to be re-used immediately. After regeneration, plugs can also be stored, without their end caps, in binding buffer A or in 0.1% sodium azide (made up in distilled water) at 2-8°C until further use.





**Used Spin Column** 



# Easy-to-read Midi Protocol E.g. Human serum

Fraction	Volume	Step	RCF	Spin Time	
Pre-equilibration	10 ml	BBA pH 7.4	500 g	3 min	
Sample Loading	20 ml	1:1 serum: BBA pH 7.4	150 g	30 mtn	
Wash #1	10 ml	BBA pH 7.4	500 g	3 min	
Wash #2	10 ml	BBA pH 7.4	500 g	3 min	
Final Eluate #1	10 ml	EB2 → 1.3 ml NBC	500 g	3 min	
Final Eluate #2	10 ml	EB2 → 1.3 ml NBC	500 g	3 min	

# Easy-to-read Regeneration Midi Protocol

Fraction	Volume	Step	RCF	Spin Time
Clean-up	10 ml	EB2 pH 2.5	500 g	3 min
Wash	10 ml	BBA pH 7.4	500 g	3 min

By using the Beer-Lambert law,  $A = \tilde{\epsilon} c.l$ , the concentration of lgG (mg/ml) in the sample can be measured by multiplying the absorbance at 280 nm by 0.72. These antibody concentrations are only estimates as other contaminating proteins can also contribute to the absorbance reading. However, they can provide a reliable and quantitative method for determining the concentrations of pure antibody solutions. Most researchers use a sandwich ELISA assay to accurately measure antibody concentrations within a range of 1 mg/ml to 20 mg/ml sample.

The antibodies can also be monitored for purity by SDS-PAGE under reducing or non-reducing conditions. Note that IgG appears in a reducing SDS-PAGE as 25 kDa and 50-55 kDa bands (Fig. 2, page 33). Recovery of immunoglobulins can be quantified by a standard protein Assay, scanning

densitometry of reducing or non-reducing SDS-polyacrylamide gels or ELISA. Antigen binding parameters can be measured for both affinity and avidity.

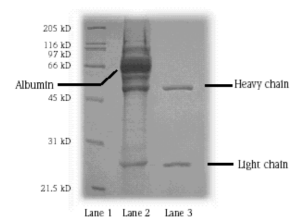


Fig. 3. Reducing SDS-Polyacrylamide gel of mouse IgG1 purified from cell culture supernatant (+ foetal calf serum) with the UptiSpin Protein G kit

Lane i Lane 2 Lane 3

Lane 1: Molecular weight markers.

Lane 2: Hybridoma cell culture supernatant before loading on to the UptiSpin column.

Lane 3: Pure mouse IgG1 eluted from the UptiSpin Protein A spin column

Fig. 4a. Graph showing that typical IgG capacity of a Protein A **Mini** spin column remains unaffected after 3 re-uses using rabbit serum. All data is within 5 % of mean recovery (1.32ma) and the % CV is 0.28 %.

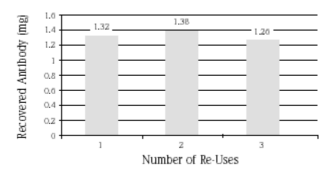
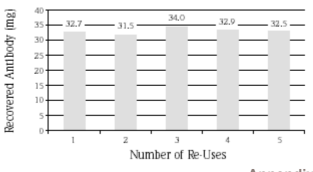


Fig. 4b. Graph showing that typical IgG capacity of a Protein A **Midi** spin column remains unaffected after 5 re-uses using rabbit serum. All data is within 5 % of mean recovery (32.7ma) and the % CV is 2.5 %.



**Appendix** 

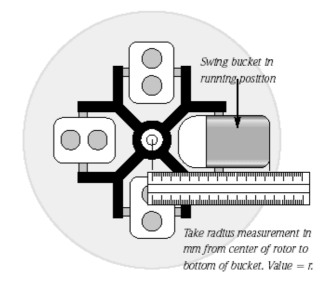
# How to convert rpm to g force (RCF) using a swing bucket rotor

It is important that the UptiSpin columns are centrifuged at the correct speeds. Use of considerable higher speeds than those indicated may damage the resin matrix and result in reduced performance. Many centrifuges display only rpm.

See the diagram to enable accurate conversion between rpm and RCF (g force). This formula will work on any rotor providing an accurate measurement is taken from the center of the rotor to the bottom of the swing bucket at its open position (when the bucket is rotated through 90° in its running position).

$$RCF = 1.12 \times r \left(\frac{rpm}{1000}\right)^2$$

Eg. 500 g corresponds to 1,670 rpm when the radius (r) = 160 mm  $\,$ 



## How to convert rpm to g force (RCF) using a 45° fixed angle rotor

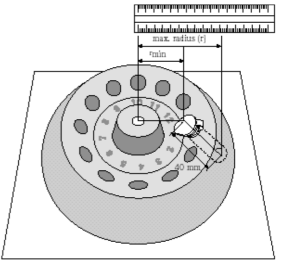
It is important that the UptiSpin columns are centrifuged at the correct speeds. Use of considerable higher speeds than those indicated may damage the resin matrix and result in reduced performance. Many centrifuges display only rpm.

See the diagram to enable accurate conversion between rpm and RCF (g force). This formula will work on any rotor providing an accurate measurement is taken from the center of the rotor to the centre of the microfuge tube lid.

$$RCF = 1.12 \times r \left(\frac{rom}{1000}\right)^{2}$$

Eg. 640 g corresponds to 3,000 rpm when the max radius (r) = 63.2 mm. (Eg. MSE Micro Centaur microfuge). 1770 g corresponds to 5,000 rpm when the max radius

Take radius measurement in mm from center of rotor to center of microcentrifuge tube lid. Value =  $r_{min}$ .



#### Determination of the maximum radius (r)

 $r = minimum radius in mm (r_{min}) + 40 mm$  (the length of the micro-centrifuge tube),  $sin 45^{\circ}$ 

#### Protein A kit buffer formulation:

Use the following recipes to prepare the buffers supplied with the UptiSpin Protein A kit. All buffers contain 0.1 % sodium azide as a preservative and can be stored at room temperature:

### Binding Buffer A (1.5 M Glycine/NaOH buffer, 3 M NaCl, pH 9.0)

Add 112.6 g glycine (free base; Mr 75.07), 175.3 g NaCl (Mr 58.44), 1.0 g NaN3 to 800 ml distilled water. Titrate with 5 M NaOH to pH 9.0. Make up final volume to 1 L with distilled water.

## Elution Buffer B1 (0.1 M Sodium citrate buffer pH 5.5)

Add 23.44 g citric acid (trisodium salt, dihydrate; Mr 294.1), 3.872 g citric acid (anhydrous; Mr 192.1), 1.0 g NaN3 to 900 ml distilled water. Make up final volume to 1 l with distilled water.

# Elution Buffer B2 (0.2 M Glycine/HCI buffer pH 2.5)

Add 15.0 g glycine (free base; Mr 75.07), 1.0 g NaN3 to 900 ml distilled water. Titrate with 5 M HCl to pH 2.5. Make up final volume to 1 l with distilled water.

# Neutralization Buffer C (1 M Tris/HCl buffer pH 9.0)

Add 103.72 g Tris base (Mr 121.1), 22.72 g Tris hydrochloride (Mr 157.6), 1.0 g NaN3 to 800 ml distilled water. Make up final volume to 1 l with distilled water.

### **Questions and Answers:**

1. What is the preferred rotor for the UptiSpin Mini and Midi spin columns?

**Mini spin columns:** The preferred rotor is a fixed angle rotor. There is no need to orientate the Mini spin column in the fixed angle rotor.

**Midi spin columns:** The preferred rotor is a swing bucket rotors. For optimal performance with a fixed angle rotor, ensure that the orientation of the spin column in the rotor is the same for sample binding, washing and elution steps.

## 2. Do I need to filter the buffers prepared in my laboratory?

It is good laboratory practice to filter all buffers. However, buffers supplied with the kit are pre-filtered for immediate use.

## 3. Do I need to pre-filter my sample before loading it on to a UptiSpin column?

All samples must be filtered through a 0.2 µm pre-filter **immediately** before loading the samples on to the spin column.

# 4. What are the typical binding capacities of UptiSpin Mini and Midi spin columns?

**Mini spin columns:** Protein A and G resin plugs have typical capacities of 1 mg rabbit IgG from serum. **Midi spin columns:** Protein A and G resin plugs have minimum capacities of 20 mg rabbit IgG from serum.



5. How should I prepare my sample for the UptiSpin column?

Many chromatographic procedures demand that the sample is pre-conditioned prior to loading. Protein A affinity separations usually require the sample to be diluted 1:1(v/v) in 1 x binding buffer. We recommend that all samples are diluted 1:1(v/v) in the binding buffer supplied with the Protein A or Protein G kit.

6. How can I process a large volume sample?

The Mini and Midi spin columns have a finite maximum volume capacity. If you have a volume of sample (>200 ml), we recommend that you use the established procedure of ammonium sulphate precipitation to concentrate your target antibody. Although many IgGs ( $\gamma$ -globulins) precipitate at a lower concentration of ammonium sulphate than most other proteins, 50% ammonium sulphate is sufficient.

7. What is the maximum volume of solution I can load on to a Mini or Midi spin column?

Mini spin columns: You can load a maximum volume of 0.65 ml.

Midi spin columns: You can load up to 20 ml in a swing bucket rotor and up to 10 ml in a fixed angle rotor.

8. What is the highest speed that I can spin the UptiSpin Mini and Midi spin columns?

**Mini spin columns:** Although the spin columns have been tested at 11,960 g (13,000 rpm in a fixed angle rotor with an average radius of 49 mm), we do not recommend spin speeds greater than 5,000 g. At very high speed, you may observe gel shrinkage away from the side walls. This will not affect the performance of the spin columns as the gel will rehydrate rapidly in subsequent spin steps.

**Midi spin columns:** There is no need to spin the devices at speeds greater than 1,250 g. No performance data is available at centrifugal speeds greater than 1,500 g.

9. Is there a minimum spin speed for the Mini and Midi spin columns?

There is no minimum spin speed for either then Mini or Midi spin columns. The devices can be spun at speeds as low as 50 g.

10. What are the minimum elution volumes from the UptiSpin spin columns?

Mini spin columns: The minimum elution volume is 0.5 ml.

Midi spin columns: The minimum elution volume is 5 ml.

11. How many times can I re-use the UptiSpin Mini and Midi spin columns?

**Mini spin columns:** Assuming that all samples are correctly filtered, sufficient buffer is provided in the kit for 3 re-uses of each Mini spin column.

**Midi spin columns:** Each Midi plug can be re-used typically up to 5 times. There is sufficient buffer volume in the kit for one complete use of each Midi spin column.

12. How can I regenerate the UptiSpin Protein A plug?

**Mini spin columns:** We recommend that you wash the plugs with 2 x 0.65 ml elution buffer B2 (pH 2.5) by centrifuging the spin columns at 1,800 g for 2 min. Then, re-equilibrate the plugs with 2 x 0.65 ml binding buffer A by centrifuging the spin columns at 1,800 g for 2 min. Proceed to the preequilibration step if plugs are to be re-used immediately. Do note that spin times of used plugs may be longer. After regeneration, plugs can also be stored, without their end caps, in a beaker containing 0.1 % sodium azide (made up in distilled water) at 2-8 °C until further use. **Midi spin columns:** We recommend that you wash the plugs with 10 ml elution buffer B2 (pH 2.5) by centrifuging the spin columns at 500 g for 3 min. Then, re-equilibrate the plugs with 10 ml binding buffer A by centrifuging the spin columns at 500 g for 3 min. Proceed to the pre-equilibration step if plugs are to be re-used immediately. Do note that spin times of used plugs may be longer. After regeneration, plugs can also be stored, without their end caps, in a beaker containing 0.1 % sodium azide (made up in distilled water) at 2-8 °C until further use.

13. Do I need to perform regeneration of the spin column immediately after the elution step?

We do not recommend storing the resin columns in elution buffer pH 2.5. Prolonged storage in pH 2.5 buffer could harm the Protein A ligand. After the elution step, continue to the regeneration procedure. The resin plugs can then be stored for re-use.

14. Can I autoclave the UptiSpin Protein A and Protein G plugs?

The UptiSpin Protein A or G plugs cannot be autoclaved.

- 15. If I experience significant fouling of the resin plug, do you recommend any cleaning-in-place procedures? Most suitable cleaning procedures tend to be determined empirically. The chosen procedure depends largely upon the nature of the previous sample loaded on the spin column.
- 16. What shall I do if the binding buffer A is translucent yellow?

This is a property of the Glycine component of binding buffer A. There is no adverse affect on the performance of the plug or the integrity of your antibody. Continue to use this binding buffer bottle for your purification.

17. Should I be concerned if the plugs partially dry out during the centrifugal steps?

The plugs are robust. Partially dried plugs rehydrate rapidly. There are no adverse effect upon the performance of the plugs.



18. Do I need to be careful with the type of chaotropic ion I use when eluting antibodies from affinity columns? It is recommended to use the mildest chaotropic agents at the lowest possible concentration that will ensure rapid elution and high recovery of activity. Iodination reactions employing either chloramine T or IODO-GEN (1,3,4,6-tetrachloro-3-6-diphenylglycouril) are particularly sensitive to inhibition by low concentrations of thiocyanate ions and, if antibodies are eluted from affinity columns or Protein A affinity columns, using this chaotropic ion, it is essential that they are dialyzed thoroughly after elution to remove thiocyanate ions.

## 19. How do I monitor purity of the isolated antibodies?

Purity is best measured by gel electrophoresis. When analyzed by SDS-PAGE under non-reducing conditions, IgG antibodies should give a single protein band of about 160-170 kDa. On reduction with DTT or 2-mercaptoethanol, two or more bands will be seen corresponding to the individual heavy chains (50-55 kDa) or light chains (25-30 kDa). Other protein bands that are visible only on reduction may point to proteolytic action. This can often be prevented by careful use of protease inhibitors in culture supernatants before storage.

- 20. What are the critical starting conditions for Protein A and G UptiSpin spin columns? Sample pH and salt concentration are usually not critical, except that the pH should be equal to or above 5.0. Protein A or G spin columns can also serve as tools for rapid buffer exchange. However, the salt concentration is normally quite high (> 0.1 M) to prevent non-specific binding.
- 21. Do I need to control the salt concentrations during Protein A and G chromatography?
  Use 0.1-0.5 M salt to reduce non-specific adsorption. When working with Protein A, use high salt (2-3 M NaCl) with high pH to promote the binding of mouse IgG1.
- 22. Is pH an important parameter to control during Protein A and G chromatography? The elution pH is the most critical variable. Protein G usually requires more acidic pH conditions to desorb the target immunoglobulins. For Protein A, elution by pH steps (starting at pH 6) may fractionate different species (weaker binding bovine IgG from target antibodies) or subclasses. High pH (pH 8-9), in conjunction with high salt may promote binding of mouse IgG1 to Protein A. The binding buffer pH should normally be higher than pH 6.0-7.0.
- 23. Can I elute antibodies from a Protein A and G spin column using divalent cations? Concentrations of divalent cations (particularly Mg<sup>2+</sup>) up to 1 M can sometimes replace acidic pH if there is concern about loss of activity of acid-labile immunoglobulins.

### Troubleshooting assistant:

Bubbles or cracks appear in the resin bed

• The spin column has been stored at a cool temperature and then rapidly warmed up. UptiSpin column should be warmed slowly to room temperature before use.

The sample does not flow easily through the spin column

- The resin is clogged with particulates. Pre-filter the sample just before loading it on to the UptiSpin column. Ascites must be delipidated before use. Please refer to page 9 for the recommended delipidation procedure.
- If the spin column are not stored at 2-8°C, or they have been used more than once and stored in the absence of a bacteriostat, microbial growth in the column may restrict flow through the resin plug.
- The centrifugal speed for the sample loading step can be increased to a maximum 1,500 g.

No elution of the target protein is observed from the spin column

- The pHof the elution buffer may be incorrect. It is advisable to prepare new solutions.
- The elution conditions are too mild to desorb the target protein.

The recovery of target protein is low

• The binding of antibodies to Protein A or G is attributed in part to hydrophobic forces. Use chaotropic salts to reduce the strength of all hydrophobic interactions.

Poor resolution of the target protein

- The sample volume or concentration may be too large for the capacity of the resin plug. In this case, reduce the sample load or sample volume.
- The sample may also need to be filtered carefully.

The target protein elutes at an unexpected position

- There may be an ionic interaction between the protein And UptiSpin resin. You should maintain the ionic strength above 50 mM.
- There may be hydrophobic interactions between the sample and UptiSpin resin. In this instance, reduce the salt concentration and add suitable detergents or organic solvents.
- The column may be dirty. All claims made of UptiSpin column are guaranteed for the first bind-wash-elute cycle only.



The elution profile cannot be reproduced

- The nature of the sample may have altered and so it may be important to prepare a fresh sample.
- The sample load may be different from the original sample load. It is advisable to keep all these parameters constant.
- Proteins or lipids may have precipitated in the resin bed. Use elution conditions which stabilize the sample.
- The buffer pHand ionic strength is incorrect and new buffers will need to be prepared.

#### Glossary:

affinity chromatography - chromatographic separation based on a specific interaction between an immobilized ligand and a binding site on a macromolecule.

antibody - an immunoglobulin produced by the immune system of vertebrates in response to exposure to a foreign substance.

antigen - a molecule which can bind specifically to an antibody.

antiserum - the serum fraction from an animal that has been immunized or exposed to an immunogen and contains antibodies to a particular antigen.

ascites - a liquid tumour formed by injection of a hybridoma cell line into the peritoneal cavity. It is a common source of monoclonal antibodies from mice.

bed volume - the total volume occupied by the chromatographic packed bed. It is also referred to as the column volume or CV.

cell culture supernatant - the fluid made during cell culture (either roller bottle, suspension or perfusion) containing tissue media components and the secreted target.

chaotropic agent - a molecule which interferes with hydro-phobic interactions by disrupting the ordered structure of water molecules. Examples include urea and quanidine.

*hybridoma* - a hybrid cell line produced by fusing antibody producing cells with myeloma cells to generate immortal reproducing cells that produce specific monoclonal antibodies indefinitely in cell culture.

immobilized - bound to a surface, usually through covalent linkages.

immunoglobulin (Ig) - comprising 5 distinct classes in most higher animals. Classes called IgG (the most common), IgM, IgA, IgD and IgE. They differ from each other in size, charge, amno acid composition and carbohydrate content.

ion exchange chromatography - chromatographic separation based on different charge properties of macromolecules. isoelectric point - the pHat which the protein has no net charge.

monoclonal antibody - an antibody derived from a single clone of immune cells. They are usually formed from a hybridoma cell line.

polyclonal antibodies - antibodies produced to the same immunogen by different cell types. Antibodies from antiserum are almost always polyclonal.

protein A/protein A - cell wall proteins of certain pathogenic bacteria which specifically bind to the Fc region of immunoglobulins.

#### References:

**Affinity Separations: A Practical Approach** 

1997 (Matejtschuk, P Ed.) IRL PRESS at Oxford University Press. ISBN: 0-19-963550-1.

Biochemistry, 2nd Edition

1996 (Mathews, CK & van Holde, KE) Benjamn/Cummngs Publ. Co. ISBN: 0-8053-3931-0.

**Purification Tools for Monoclonal Antibodies** 

1996 (P Gagnon) Validated Biosystems, USA, ISBN: 0-91.33515-9-9.

Biochemistry, 4th Edition

1995 (L Stryer) WH Freeman and Co. ISBN: 0-7167-2009-4.

Immunology

1985 (I Roitt, J Brostoff, D Male) Gower Medical Publ. Ltd., ISBN: 0-443-02912-1.

**Protein Purification Applications : A Practical Approach** 

1995 (E.L.V. Harris and S. Angal Eds.) IRL PRESS at Oxford University Press. ISBN: 0-19-963023-2.

**Basic Proteins and Peptide Protocols** 

1994 (J Walker Ed.) Humana Press, ISBN: 0-89603-269-8.

**Recombinant Protein Protocols** 

1997 (RS Tuan Ed.) Humana Press, ISBN: 0-89603-400-3.

## **Ordering information**

Cat.# Description

MINI

UPBB9720 Protein A Mini Sample Kit : 2 A Plugs (includes buffers)
UPBB9730 Protein A Mini Kit – 16 A Plugs (includes buffers)

UPBB9740 Protein A Mini Bulk Pack – 48 A plugs (does not include buffers)

MIDI

UPBB9750 Protein A Midi Kit – 4 Midi A Plugs (includes buffers)

UPBB9760 Protein A Midi Bulk Pack – 12 A Plugs (*does not include buffers*)

