

Protein A IgG Purification Kit

44667

0528.2

Number	Description
44667	Protein A IgG Purification Kit , sufficient materials to perform 45 purifications Kit Contents: Protein A Columns , 5 × 1mL resin-packed columns Support: Crosslinked, 6% beaded agarose resin Supplied Format: Packed resin in 0.02% sodium azide storage solution Binding Capacity: 6-8mg mouse IgG per milliliter of settled resin Protein A IgG Binding Buffer , 1L, pH 8.0, contains EDTA as a preservative IgG Elution Buffer , 500mL, pH 2.8, contains amine Desalting Columns , 5 × 5mL

Storage: Upon receipt store at 4°C. Kit is shipped at ambient temperature.

Introduction

The Thermo Scientific Protein A IgG Purification Kit is useful for routine purification of antibodies. Samples containing IgG are incubated with immobilized Protein A in a buffer that facilitates binding. After non-IgG components are washed from the column, the bound IgG is recovered by elution. The supplied buffers provide for maximum immunoglobulin binding and elution efficiency with the Protein A column. The pre-packed Protein A columns and desalting columns supplied in the kit make IgG purification easy and efficient.

Protein A is a cell wall component produced by several strains of *Staphylococcus aureus* and consists of a single 42 kDa polypeptide chain and contains little or no carbohydrate.^{1,2} The unique functional property of Protein A is its ability to bind specifically to the Fc region of immunoglobulins, especially IgG. The Protein A molecule contains four high affinity ($K_a = 10^8 \text{ M}^{-1}$) binding sites capable of interacting with the Fc region of several species.³ The molecule is heat-stable and retains its native conformation when exposed to denaturing agents such as 4M urea, 4M thiocyanate and 6M guanidine hydrochloride.⁴

Protein A covalently coupled to agarose is used as an affinity support for isolating IgG from several species.^{5,6,7} However, the interaction between Protein A and IgG is not equivalent for all species.^{8,9} Even within a species, Protein A interacts with some subclasses of IgG and not others. For example, human IgG₁, IgG₂, and IgG₄ bind strongly but IgG₃ does not³ and mouse IgG₁ binds poorly to Protein A.¹⁰ Despite its variable binding characteristics, Protein A possesses IgG-binding properties that make it ideal for affinity purification of IgG from the serum. When using samples from species not well referenced in the literature, conduct preliminary experiments to determine if a particular immunoglobulin binds.

The Protein A resin supplied in this kit is superior to other commercially available Protein A agarose resins because immobilization is accomplished by stable secondary amine bonds. The result is an affinity resin with excellent stability and binding characteristics, enabling the column to be re-used multiple times without significant loss in binding capacity.

Important Product Information

- The Protein A Binding Buffer supplied provides the highest efficiency of IgG binding and elution for most species. Using other buffer formulations may significantly alter the binding capacity and the wash volumes required for efficient purification. Therefore, optimization may be necessary when using other buffers.
- For optimal recovery, use a sample size such that the expected IgG load on the column is less than 80% of the maximum binding capacity. The total IgG content of serum is approximately 10-15mg/mL. The concentration of antibody in tissue culture supernatant varies considerably among hybridoma clones. Be aware that antibodies from fetal bovine serum (FBS) culture media supplement will be purified along with the antibody of interest.
- The crosslinked 6% beaded agarose resin can tolerate commonly used water-miscible solvents when they are added in a stepwise gradual manner. The agarose resin will compress under pressure causing column flows to slow. Never freeze agarose resin, as this will cause irreversible damage to the bead structure.
- Serum samples, ascites fluid, plasma or tissue culture supernatant may be used with this product.

Reagent Preparation

Kit components	Equilibrate Immobilized Protein A and all buffers to room temperature.
Neutralization Buffer	Prepare 1mL of high-ionic strength alkaline buffer such as 1M phosphate or 1M Tris; pH 7.5-9
Antibody Sample	To ensure proper ionic strength and pH for optimal binding, dilute serum, ascites fluid or tissue culture supernatant samples with an equal volume of Binding Buffer. Centrifuge cloudy samples and use only the clear supernatant. To obtain good IgG recovery from lipemic plasma or serum, centrifuge diluted samples at 10,000 × g for several minutes.

Procedure for Purification of IgG

A. Equilibrate Protein A Column

1. Sequentially remove top and bottom cap from Protein A column.
Note: Always remove top cap first and bottom cap second to prevent air bubbles from being drawn into the gel.
2. Pour off and discard the storage solution, which contains 0.02% sodium azide.
3. Equilibrate Protein A column by applying 3-5mL of Binding Buffer and allowing it to drain through the column.
Note: In all steps column flow will stop when solution drains down to the disc at the top of the gel bed. This prevents the gel bed from drying out. However, do not leave drained column uncapped for more than a few minutes.

B. Apply Sample to Column

1. Apply up to 5mL of Antibody Sample to the equilibrated Protein A column.
Note: With dilute antibody samples (for which 10-50mL of sample is needed to approach the column's binding capacity), better yield can be obtained more quickly by incubating the sample and immobilized Protein A in batch format. Remove the top disc from the column and use a portion of the sample to wash the resin from column into a tube that is large enough to hold the entire volume of sample and resin. Incubate the resin-sample slurry for 1-2 hours at room temperature or overnight at 4°C with gentle mixing. Pour the slurry back into the column to re-pack the resin bed before proceeding to wash and elution steps.
2. Allow Antibody Sample to flow through the column.
Note: If desired, collect the solution in a clean collection tube and reserve it for binding efficiency analysis. If Antibody Sample contains more IgG than can bind to the Protein A column (or is an antibody type that does not bind to Protein A), the collected flow-through will contain the excess antibody. By saving the flow-through, non-bound antibody can be recovered and examined by antibody-specific assays.

3. Wash column by adding 5-15mL of Binding Buffer and allowing it to drain through the column.

Note: If desired, verify that all unbound proteins are thoroughly washed away by collecting separate 2mL fractions of the solution as it drains and measuring their absorbance at 280 nm. Complete washing is indicated by fractions having absorbances similar to Binding Buffer alone.

C. Elute Antibody

1. Elute the bound IgG with 5-10mL of the IgG Elution Buffer, collecting separate 1mL fractions of the eluate that drains from the column.

Note: If desired, each 1mL fraction can be neutralized by adding 50 μ L of 1M Tris, pH 9.5 or 100 μ L of Binding Buffer. Neutralization helps to stabilize antibodies that may otherwise be inactivated by long-term storage in acidic buffer.

2. Determine which fractions contain antibody by measuring the absorbance of each one at 280 nm. Pool fractions having the highest absorbance values (typically fractions 2, 3 and 4) compared to neutralized Elution Buffer alone.
3. The purified antibody in neutralized Elution Buffer suitable for storage, Thermo Scientific Pierce BCA Protein Assay (Product No. 23225), SDS-PAGE or dilution for immunoassays. Alternatively the buffer may be exchanged to a system compatible with the specific downstream application (see optional procedure that follows).
4. Regenerate the Immobilized Protein A column by washing with 5-10mL of Elution Buffer. Columns may be regenerated a minimum of 10 times without significant loss of binding capacity.
5. For storage, wash column with 5mL of water containing 0.02% sodium azide. When approximately 2mL of solution remains above the top disc, replace the bottom cap on the column followed by the top cap. Store column upright at 4°C.

Exchange Eluted Antibody into Different Storage Buffer (Optional)

A. Alternative Methods for Buffer Exchange

In the purification procedure, IgG is eluted from Protein A using low-pH, amine-containing buffer and then neutralized with Tris or phosphate buffer. Often, this neutralized buffer is suitable for long-term storage of the antibody; however, certain applications will require a buffer exchange. For example, most biotinylation or enzyme-labeling procedures will necessitate removing amines from the buffer.

Two options exist for buffer exchange. One is to dialyze the purified antibody against the buffer of choice. For this purpose, choose a dialysis cassette (see Related Thermo Scientific Products) appropriate for the volume of purified antibody solution. Use Thermo Scientific Slide-A-Lyzer Concentrating Solution to concentrate the antibody solution after dialysis.

The second option for buffer exchange is size-exclusion chromatography or gel filtration. For this purpose, the kit includes desalting columns. The porous resin beads have an average molecular weight exclusion limit of 5,000. Therefore, molecules with molecular mass greater than 5kDa will pass by the resin beads and emerge in the column void volume, while smaller molecules will pass through the column more slowly as they migrate into and through the resin. In this case, the purified IgG (approximately 150kDa) will emerge from the column first and be separated from the buffer salts that will emerge later. To achieve adequate separation, apply less than 1.25mL of sample to the 5mL desalting column. As such, two or three separate columns may have to be used to process the entire volume of purified IgG sample.

The choice between dialysis and desalting is a matter of speed vs. effort. Dialysis requires several hours to overnight to perform, but the entire sample can be processed at once with minimum hands-on time. Desalting can be performed in much less time but requires careful sample collection and identification of fractions containing the molecule of interest.

B. Procedure for Buffer Exchange Using Desalting Columns

1. Prepare 100mL of desalting buffer (i.e., the buffer into which the purified antibody will be exchanged).
2. Equilibrate desalting columns to room temperature. Use a separate column for each 1.25mL of antibody sample.
3. Invert column several times to resuspend the resin. Position the column upright in a test tube or clamp and allow the resin to settle for several minutes.
4. Remove the top cap from the column and carefully pipette the storage solution (contains 0.02% azide) until 5-10mm of solution remains above the resin bed.

5. (Optional) Using the open end of the supplied resin separator, insert and slide a porous disc to within 1mm of the resin bed. A top porous disc provides a stop-flow function that prevents disturbance and drying of the resin during use.
6. Twist off column bottom end tab.
7. Equilibrate the column by adding five resin-bed volumes of buffer to the column and allowing it to drain through.
Note: The gravity-flow rate of column is approximately 0.1mL per minute. Column flow will stop when solution drains down to the disc at the top resin; however, do not leave drained column uncapped for more than a few minutes.
8. Number 6-10 collection tubes.
9. Apply up to 1.25mL of antibody sample to column and immediately begin to collect a 1mL fraction into the first numbered collection tube.
10. When column flow stops as a result of solution draining down to the top disc, add desalting buffer and continue collecting the solution as it drains. After the first 1mL fraction (#1), begin to collect 0.5mL fractions into successively numbered collection tubes until at least five 0.5mL fractions (#2-6) have been collected.
11. Measure the absorbance at 280nm of each 0.5mL fraction relative to the desalting buffer. Fractions #2-4 will likely contain the purified antibody (now exchanged into desalting buffer) as indicated by their higher absorbance values. These fractions may be pooled.
Note: If fraction #6 has absorbance significantly greater than desalting buffer alone, go back to step 7 and collect additional fractions until the absorbance approaches that of the desalting buffer. Molecules smaller than the resin's exclusion limit (i.e., the buffer salts in the starting antibody sample) will emerge from the column in subsequent fractions (e.g., fractions #8-10). If collected, these fractions may be discarded after confirming that all fractions containing antibody have been identified and saved.
12. The desalting columns can be regenerated for use again by washing with an additional 15mL of desalting buffer after the antibody has been collected. For storage, include 0.02% sodium azide in the buffer used to regenerate the column. Cap column when approximately 2mL of solution remains above the resin. Securely cap top of column and store at 4°C.

Troubleshooting

Problem	Possible Cause	Solution
Protein A column gravity-flow is exceedingly slow (i.e., < 0.1mL/minute)	Outgassing of buffers or sample on the column, has blocked resin pores with microscopic air bubbles	Degas buffers and remove air bubbles from column (see Tech Tip in Additional Information section)
No protein detected in any elution fractions with absorbance at 280nm or general protein staining of electrophoresed sample	Accidentally used desalting column instead of Protein A column for purification procedure	Use correct column in kit
	Sample devoid of any antibody species or subclass that binds to Protein A	Ensure by ELISA or isotyping kit, that the sample contains target IgG (see Related Thermo Scientific Products)
	Antibody bound to Protein A column (determined by depletion from the starting sample) but did not elute with the low-pH IgG Elution Buffer	Try Gentle Ag/Ab Elution Buffer (see Related Thermo Scientific Products)
Considerable antibody purified, but no specific antibody of interest detected	Antibody of interest is at very low concentration or has very low binding affinity for Protein A relative to other immunoglobulins in the sample	Use serum-free medium for cell supernatant samples
		Affinity purify the antibody using the specific antigen coupled to an affinity support (see Related Thermo Scientific Products)
Antibody was purified, but it is degraded (determined by lack of function in downstream assay)	Antibody is sensitive to the low-pH IgG Elution Buffer	Try Gentle Ag/Ab Elution Buffer (see Related Thermo Scientific Products)
	Downstream application is sensitive to neutralized IgG Elution Buffer	Desalt or dialyze eluted sample into suitable buffer

Additional Information

Please visit the website for the following related items:

- Tech Tip #27: Optimize elution conditions for immunoaffinity purification
- Tech Tip #7: Remove air bubbles from columns
- Tech Tip #29: Degas solutions for use in affinity columns
- Tech Tip #34: Binding characteristics of Protein A, G, A/G and L for immunoglobulins

Related Thermo Scientific Products

21001	Protein A IgG Binding Buffer, 1L
21004	IgG Elution Buffer, 1L
21027	Gentle Ag/Ab Elution Buffer, 500mL
89948-89951	NAb™ Spin Kits for Antibody Purification (Protein A, G, A/G and L, respectively)
89978-89881	NAb Spin Kits for Antibody Purification (Protein A, G, A/G and L, respectively)
20333	Immobilized Protein A, 5mL
45212	Melon™ Gel IgG Purification Kit, alternative purification method for antibody purification
44894	AminoLink® Plus Immobilization Kit, covalent attachment of proteins for affinity purification
37501	Monoclonal Antibody Isotyping Kit I (HRP/ABTS)
66382	Slide-A-Lyzer® Dialysis Cassette Kit, 10 dialysis cassettes, each appropriate for 0.5-3.0mL samples
66528	Slide-A-Lyzer Concentrating Solution, for use with 0.5 to 3mL cassettes, 200mL
66529	Slide-A-Lyzer Concentrating Solution, for use with 3 to 15mL cassettes, 500mL

References

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