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F(ab')2 Micro Preparation Kit

**BIOK2025** 

F(ab')2 Micro Preparation Kit, contains sufficient reagents to generate and purify F(ab')2

fragments from up to ten 0.125mL samples containing 25-250µg of IgG

**Kit Contents:** 

**Immobilized Pepsin** 0.5mL\*1

1.5mL\*10 Microcentrifuge Tubes

Protein A Column 0.2mL\*2

Phosphate Buffered Saline 1 pack for 1L

30mL\*1 IgG Elution Buffer pH 2.8

Microcentrifuge Tubes 2.0mL\*30

30mL\*1 F(ab')2 Digestion Buffer pH 4.4

**Desalting Columns** 0.5mL\*10

**Storage:** Upon receipt store kit at 4-8°C. Kit is shipped at ambient temperature

The F(ab')2 Preparation Kit enables efficient generation of F(ab')2 from IgG. This kit uses

Immobilized Pepsin, a nonspecific endopeptidase that is active only at acid pH and irreversibly

denatured at neutral or alkaline pH. Pepsin digestion typically produces a F(ab')2 fragment

(~110kDa by SDS-PAGE under non-reducing conditions) and numerous small peptides of the Fc

portion (Figure 1). The resulting F(ab')2 fragment is composed of a pair of Fab' units connected

by two disulfide bonds. The Fc fragment is extensively degraded and can be separated from

F(ab')2 by dialysis, gel filtration or ion exchange chromatography.

This kit contains the necessary components for F(ab')2 generation and subsequent purification.

Immobilized Pepsin is advantageous because the digestion can be immediately stopped by simply

removing the resin from the antibody digest solution. The included Spin Columns allow easy

manipulation of the resin and maximum F(ab')2 recovery. The prepacked, immobilized Protein A



Column binds the large Fc fragments and undigested IgG, allowing the F(ab')2 fragments to pass through the column for efficient purification. This complete kit makes F(ab')2 generation and purification simple, fast and effective.

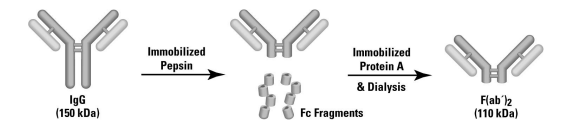


Figure 1. Schematic for preparing F(ab')<sub>2</sub> using Immobilized Pepsin and Protein A

### **Important Product Information**

- The kit components and protocol is for 125μL samples containing 25-250μg of IgG per sample.
- Proper sample preparation is essential for successful fragment generation using this kit.
   If the IgG sample contains a carrier protein such as BSA, removing it before performing the buffer exchange.
- For best results, use rabbit, human or mouse IgG. Fragmentation of IgG from other species may require optimization. For purification, the IgG species must be able to bind to Protein A.
- Digestion effectiveness will vary depending on antibody preparation and source (rate
  and completeness of digestion: rabbit > human > mouse ≥ goat). Digestion times in the
  protocol result in > 90% digestion of IgG using serum purified by Protein A or G affinity
  chromatography. Digestion over 3 hours is not recommended.

#### **Additional Materials Required**

Incubator capable of maintaining 37°C

Microcentrifuge capable of  $5000 \times q$ 

Variable speed centrifuge

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1.5mL microcentrifuge tubes

End-over-end mixer or tabletop rocker

**Material Preparation** 

Phosphate-buffered Saline (PBS): Dissolve contents of a package in 1L of ultrapure water. For

long-term storage, add 0.05% sodium azide and store at 4°C.

Procedure for Generating and Purifying F(ab')2 Fragments

A. Immobilized Pepsin Equilibration

1. Gently swirl the Immobilized Pepsin vial to obtain an even suspension. Seat the

spin column frit with an inverted 200µL pipette tip.

2. Tplace 65µL of the 50% slurry (i.e., 32.5µL of settled resin) into the 1.5mL

Microcentrifuge Tubes at  $5000 \times g$  for 1 minute and discard buffer.

3. Wash resin with 130 $\mu$ L of Digestion Buffer. Centrifuge column at 5000  $\times$  g for 1 minute

and discard buffer.

**B.** IgG Sample Preparation

1. Desalting Column, Centrifuge column at 1500  $\times$  g for 1 minute to remove storage

solution. Place a mark on the side of the column where the compacted resin is slanted

upward. Place column in centrifuge with the mark facing outward in all subsequent

centrifugation steps.

**Note:** Resin will appear compacted after centrifugation.

2. Add 300µL of Digestion Buffer to column. Centrifuge at  $1500 \times q$  for 1 minute to remove

buffer. Repeat this step three additional times, discarding buffer from the collection

tube.

3. Place column in a new Microcentrifuge Tubes, remove cap and slowly apply 125µL of

sample to the center of the compacted resin bed.

4. Centrifuge at  $1500 \times q$  for 2 minutes to collect the sample. Discard the column after use.

5. If IgG sample is 0.2-2mg/mL (i.e., 25-250μg), no further preparation is necessary. If sample

volume is less than 125μL, add Digestion Buffer to a final volume of 125μL.

C. Fragment Generation

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1. Add 125µL of the prepared IgG sample to the Microcentrifuge Tubes containing the

equilibrated Immobilized Pepsin (Section A). Briefly vortex to mix.

2. Incubate digestion reaction for 2 hours for rabbit or human IgG or 3 hours for mouse IgG

with an end-over-end mixer or tabletop rocker at 37°C. Maintain constant mixing of resin

during incubation.

3. Centrifuge at  $5000 \times q$  for 1 minute to separate digest from the Immobilized

Pepsin.

4. Wash resin with 130 $\mu$ L of PBS. centrifuge at 5000  $\times$  g for 1 minute. Repeat this step

once.

5. Add both wash fractions to the digested antibody. Total sample volume should be

385µL. Discard the Immobilized Pepsin.

Note: For best results, evaluate the digest and wash fraction via SDS-PAGE to assess

digestion completion. Protein A purification is only required to remove undigested IgG.

F(ab')2 and degraded Fc do not bind to Protein A. The resulting F(ab')2 in non-reducing

SDS-PAGE derived from human and mouse IgG will migrate with an apparent molecular

weight of ~110kDa. Rabbit F(ab')<sub>2</sub> will migrate with a lower apparent molecular weight of

~88kDa.

D. F(ab')2 Purification

1. Equilibrate the Protein A Plus Spin Column, PBS and IgG Elution Buffer to room

temperature. Set centrifuge to  $1000 \times g$ .

Place column in a collection tube and centrifuge for 1 minute to remove storage

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3. To equilibrate column, add 400µL of PBS and briefly mix. Centrifuge for 1 minute and

discard the flow-through. Repeat this step once.

4. Apply 25-500μL sample to column and cap the top tightly. Resuspend the resin and

sample by inversion. Incubate at room temperature with end-over-end mixing for 2

minutes.

5. Place column in a new collection tube and centrifuge for 1 minute. Save the

flow-through as this fraction contains F(ab')2 and Fc fragments that are too small to

bind to Protein A.

6. For optimal recovery, wash column with 200µL of PBS. Centrifuge for 1 minute and

collect flow-through. Repeat and combine wash fractions with the F(ab')<sub>2</sub> fraction from

Step 5.

7. Measure protein concentration using the BCA Protein Assay or by measuring the

absorbance at 280nm. Use an estimated extinction coefficient of 1.4. Assuming

complete IgG digestion, F(ab')2 yields may vary from 50 to 70%, depending on the

amount of starting antibody and the protein assays used.

8. If desired, perform dialysis (50K MWCO), gel filtration or ion-exchange chromatography

to remove the Fc fragments that are too small to bind to Protein A.

E. Regeneration of the Immobilized Protein A Plus Spin Column

1. Apply 400µL of IgG Elution Buffer to the Protein A Plus Spin Column. Centrifuge for 1

minute. Repeat this step two times to obtain three fractions, which will contain

undigested IgG. To save the undigested IgG, add 40µL of a neutralization buffer (e.g.,

1M phosphate or 1M Tris at pH 8-9) to each elution fraction.

2. Add 400µL of IgG Elution Buffer to the column and centrifuge for 1 minute. Discard

flow-through and repeat.

3. Add 400µL of PBS to the column and centrifuge for 1 minute. Discard flow-through and

repeat two times.



4. For storage, add  $400\mu$ L of 0.02% sodium azide in PBS to column. Replace top and bottom caps. Store column upright at 4°C. Columns can be regenerated at least 10 times without significant loss of binding capacity.

# F. Troubleshootin

Problem	Possible Cause	Solution
Low amounts of F(ab') <sub>2</sub>	lgG sample was not in	Dialyze or buffer exchange IgG into
produced as determined by	Digestion Buffer	Digestion Buffer, or decrease the
non-reducing SDS-PAGE		Digestion Buffer pH to 3-4.3 <b>Note:</b>
		Decreasing the pH might increase the
		F(ab') <sub>2</sub> amount produced but can reduce
		its immunoreactivity]
	Sample loading buffer	Use SDS loading buffer that does not
	contains reducing reagent	contain
		β-mercaptoethanol, DTT or TCEP
	Resin was not	Wash resin with 0.5mL of Digestion
	equilibrated in Digestion	Buffer before adding IgG sample
	Buffer before adding IgG	
	Sample is goat or mouse	Reduce IgG concentration and increase
	IgG <sub>1</sub>	digestion time to 8 hours
	Some mouse IgG <sub>1</sub> are	Use the IgG <sub>1</sub> Fab and F(ab') <sub>2</sub> Preparation
	resistant to pepsin cleavage <sup>1</sup>	kit
	Sample contains protein	Remove BSA with the Antibody Clean-up
	other than IgG (e.g., BSA),	Kit
	which can increase	
	digestion time	
F(ab')₂ has low immunoreactivity	Sample digested for too long	Reduce digestion time; do not exceed 8
		hours
	The low pH of Digestion	Use the IgG <sub>1</sub> Fab and F(ab') <sub>2</sub> Preparation



	Buffer decreased F(ab') <sub>2</sub>	Kit
	activity	
Low F(ab´) <sub>2</sub> recovery	Incomplete washing of the	Two 130μL washes of PBS are required
	pepsin resin	for maximum recovery
A portion of undigested IgG does	Sample is goat or mouse	Goat IgG binds weakly to Protein A, so try
not bind to Protein A	$lgG_1$	an alternative purification method such as
		ion-exchange
		Dilute sample in Protein A Binding Buffer
		before adding to the Protein A Plus Spin
		Column