

Mouse IgG1 Fab Preparation Kit

Cat No: BIOK2025-1

Size: 10 Antibody Samples

PRODUCT DESCRIPTION

Mouse IgG1 Fab and F(ab)2 Preparation Preparation Kit enables efficient Fab generation from IgG. This kit uses Ficin, a nonspecific thiol-endopeptidase, immobilized on agarose resin. Immobilized enzyme is advantageous because digestion can be immediately stopped by simply removing the IgG solution from the resin, resulting in a digest that is enzyme-free. Digestion by Ficin produces 50kDa Fab and Fc fragments.

This complete kit makes Fab generation and purification simple, fast and effective. The kit includes spin columns for easy manipulation of the enzyme resin.

PROVIDED MATERIALS

Available for: 10 antibody samples, each containing 0.25 to 4 mg of IgG

- Ficin enzyme agarose, 1.25 mL
- Cysteine hydrochloride, 1 g
- Fab digestion buffer, 120 mL
- NAb Protein A Plus centrifuge column, 1 mL, 1 column
- PBS packages (each can prepare 500 mL), 2 packages
- IgG elution buffer, 120 mL
- Zeba desalting centrifuge column, 7K MWCO, 2 mL, 10 columns
- Centrifugal column, 0.8 mL, 10 columns
- Microcentrifuge tube, 2 mL, 30 tubes

Storage method: Store at 2-8 $\,^\circ\!\mathbb{C}\,$ for more than 6 months, do not freeze.

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Important Product Information

• These instructions are optimized for rabbit, human and mouse IgG (250ug-4mg per sample) Fragmentation of IgG from otherr species might require optimization. For purification, the IgG species must be able to bind to Protein A. For mouse IgG1, use the IgG1 Fab and F(ab['])2 Preparation Kit.

- components and protocol are for 0.5mL samples containing 0.25-4mg lgG. For 25-250 μ g samples use the Fab Micro Preparation Kit.

• Proper sample preparation is essential for successful fragment generation using this kit. If the IgG sample contains a carrier protein such as BSA, use the Antibody Clean-up Kitto remove it before performing the buffer exchange.

Additional Materials Required

- Incubator capable of maintaining 37 $^\circ\,$ C
- Microcentrifuge capable of 5000 $\, imes\,$ g
- Variable speed centrifuge
- 15mL conical collection tubes
- End-over-end mixer or tabletop rocker

Material Preparation

• Digestion Buffer: Dissolve 35mg cysteine•HCl in 10mL of the supplied Fab Digestion Buffer (pH

10). After adding the cysteine • HCl the pH should be ~7.0.

Note: Cysteine readily oxidizes to cystine; therefore, prepare this buffer on the same day of use.

• Phosphate-buffered Saline (PBS): Dissolve contents of a package in 500mL of ultrapure water.

For long-term storage, add 0.05% sodium azide and store at 4 $^\circ\,$ C.

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Procedure for Fab Generation and Purification

A、 Immobilized Ficin Equilibration

1. Gently swirl the Immobilized Ficin vial to obtain an even suspension. Seat the spin-column frit with an inverted 200μ L pipette tip.

2. Twist off the bottom tab from a 0.8mL spin column and place into a 2mL microcentrifuge tube. Using a wide-bore or cut pipette tip, place 0.25mL of the 50% slurry (i.e., 0.125mL of settled resin) into the 0.8mL spin column. Centrifuge the column at 5000 \times g for 1 minute and discard buffer.

3. Wash resin with 0.5mL of Digestion Buffer. Centrifuge column at 5000 $\, imes\,$ g for 1 minute and discard buffer. Cap bottom of spin column with supplied rubber cap.

B、 IgG Sample Preparation

1. Twist off the bottom closure of a Zeba Spin Desalting Column and loosen cap. Place column in a 15mL collection tube.

2. Centrifuge column at 1000 \times g for 2 minutes 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.

3. Add 1mL of Digestion Buffer to column. Centrifuge at 1000 $\,\times\,$ g for 2 minutes to remove buffer. Repeat this step three additional times, discarding buffer from the collection tube.

C、 Generation of Fragments

1. Add 0.5mL of the prepared IgG sample to the spin column tube containing the equilibrated Immobilized Ficin. Place top cap and bottom plug on spin column.

2. Incubate the digestion reaction for the appropriate time (see Appendix A) with an end-over-end mixer or a tabletop rocker at 37° C. Maintain constant mixing of resin during incubation.

3. Remove bottom cap and place spin column into a microcentrifuge tube. Centrifuge column at 5000 \times g for 1 minute to separate digest from the Immobilized Ficin.

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4. Wash resin with 0.5mL PBS. Place spin column into a microcentrifuge tube. Centrifuge column at 5000 $\, imes\,$ g for 1 minute.

5. Add the wash fraction to the digested antibody from Step 3 Total sample volume should be 1.0mL. Discard the used Immobilized Ficin.

Note: To assess digestion completion, evaluate the digest and wash fraction via SDS-PAGE. The separated digest and wash fraction contains cysteine. Boiling samples in non-reducing SDS-PAGE loading buffer will reduce the sample. To avoid reducing the 50kDa Fab fragment on SDS-PAGE, do not boil the samples. See representative gel in Appendix B.

D、Fab Purification

1. Equilibrate the NAb Protein A Plus Spin Column, PBS and IgG Elution Buffer to room temperature. Set centrifuge to 1000 $\, imes\,$ g.

 Loosen top cap on spin column and snap off bottom closure. Place column in a 15mL collection tube and centrifuge for 1 minute to remove storage solution (contains 0.02% sodium azide).
 Discard the flow-through.

3. Equilibrate column by adding 2mL of PBS, centrifuge for 1 minute and discard the flow-through. Repeat this step once.

4. Cap bottom of column with the included rubber cap. Apply sample to column and tightly cap top. Resuspend the resin and sample by inversion. Incubate at room temperature with end-over-end mixing for 10 minutes.

5. Loosen top cap and remove bottom cap. Place column in a new 15mL collection tube and centrifuge for 1 minute. Save the flow-through as this fraction contains Fab fragments.
6. For optimal recovery, wash column with 1mL of PBS. Centrifuge for 1 minute and collect the

flow-through. Repeat and combine wash fractions with the Fab fraction from Step 5.

7. Apply 1mL of IgG Elution Buffer to the NAb Protein A Plus Spin Column and centrifuge for 1 minute. Repeat this step two times to obtain three fractions, which will contain undigested IgG and Fc fragments. To save the undigested IgG or Fc fragments, add 100μL of a neutralization buffer (e.g., 1M phosphate or 1M Tris at pH 8-9) to each of the elution fractions.
8. Measure protein concentration by absorbance at 280nm. Use an estimated extinction

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coefficient of 1.4. Assuming complete IgG digestion, Fab yields may vary from 50 to 65%, depending on the amount of starting antibody and the protein assays used. Protein concentration may also be measured using the Reducing Agent Compatible BCA Protein Assay; however, the sample must contain less than 2.5mM cysteine. The undiluted digest and Protein A fraction contains approximately 5mM cysteine.

E、Regeneration of the Immobilized Protein A Column

1. Add 3mL of IgG Elution Buffer and centrifuge for 1 minute. Repeat and discard flow-through.

2. Add 3mL of PBS to the column, centrifuge for 1 minute and discard the flow-through. Repeat three times.

3. For storage, add 3mL 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.

Troubleshooting

Problem	Possible Cause	Solution	
Low amounts of Fab	The IgG sample was not properly	Dialyze or buffer-exchange IgG	
(50kDa) produced as	prepared	into the Digestion Buffer	
visualized by	Cysteine in the Digestion Buffer	Prepare Digestion Buffer with	
nonreducing SDS-PAGE	oxidized to cystine	cysteine on the same day of	
		usage	
	Sample loading buffer contains	Use SDS loading buffer that does	
	reducing reagent	not contain β	
		-mercaptoethanol, DTT or TCEP	
	Digested material contains	Desalt before SDS-PAGE	
	cysteine		
	Resin not equilibrated in	Wash resin with 0.5mL of	
	Digestion Buffer	Digestion Buffer before adding	

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		IgG sample
	Sample contains protein other	Purify the antibody sample with
	than IgG (e.g., BSA), which can	the Pierce Antibody Clean-up Kit
	increase digestion time	
Fab has low	Sample was digested for too long	Reduce digestion time and do
immunoreactivity		not exceed 20 hours or try using
		the Pierce F(ab ´)2 Preparation
		Kit
A portion of undigested	Sample is goat IgG	Try an alternative purification
IgG or Fc does not bind		method such as ion-exchange
to Protein A		chromatography
	Sample is mouse IgG1	Dilute mouse IgG1 sample in
		Protein A Binding Buffer before
		adding to the NAb Protein A Plus
		Spin Column

Appendix

A. Recommended Digestion Times

This kit is for digesting ten 0.5mL samples of rabbit, human or mouse IgG at 0.5-8mg/mL. Digestion effectiveness will vary depending on antibody preparation and source (rate and completeness of digestion: mouse> rabbit > human). Digestion times listed in Table 1 result in > 90% digestion for mouse and rabbit IgG and > 80% digestion for human IgG. Data was generated using serum purified by Protein A or G affinity chromatography. No significant increase in digestion is obtained for more than 10 hours. Extended digestion times > 20 hours can degrade Fc, which might not bind to Protein A.

Species	<u>lgG</u> (mg/mL)	Digestion Time (hours)
Rabbit	8	8-9
	4	6-7
	1.5	4-5
	0.5	3-4
	8	5-6
Human	4	5-6
numan	1.5	3-4
	0.5	2-3
5	8	4-5
Marria	4	3-4
Mouse	1.5	2-3
	0.5	2-3

Table 1. Recommended digestion times for various species and concentrations of IgG.

B. Protein Gel Interpretation

The Fab and Fc analyzed by non-reducing and non-boiled SDS-PAGE typically migrate with an apparent molecular weight of 45-50kDa, depending on the antibody species. In reducing SDS-PAGE, Fab fragments migrate near 25kDa, and Fc fragments migrate at 28-30kDa. The presence of the Fc at 28-30kDa confirms digestion of IgG. Boiling the IgG digest before gel loading will result in a reduced sample, because of the cysteine present. Also, an additional band might be present in reduced SDS-PAGE, which is likely the undigested IgG heavy chain (50kDa).

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