

Mouse IgM Specific Enhanced Chemiluminescent Western Blot Kit - KOA0134

Mouse IgM Specific Enhanced Chemiluminescent Western Blot Kit is a four-component system for sensitive detection of membrane-immobilized proteins on western blots probed with mouse primary antibodies using HRP-conjugated anti-mouse secondary antibodies and enhanced chemiluminescent peroxidase substrate reagents.

KIT COMPONENTS

Mouse IgM Specific Enhanced Chemiluminescent Western Blot Kit					
Component	Catalog Number	Description	Size		
Blocking Reagent	KOB0104	Blotto Blocking Buffer	10 g		
Peroxidase Conjugated Secondary Antibody	KOL0305	Anti-Mouse IgM (Goat) Peroxidase Conjugated Antibody	100 μL		
Chromogenic Reagent	KOG0101A KOG0101B	Two Component Chromogenic Reagent for Enhanced Chemiluminescence (20X)			

II. STORAGE

Store at 4°C for one year. Avoid freezing.

III. REQUIRED EQUIPMENT AND REAGENTS

- APES or POLY-L-LYSINE
- Nitrocellulose or PVDF membrane.
- Diluent Buffer (pH 7.2~7.6)
 - Diluent Buffer (for preparation of blocking buffer and antibody solution):
 - Add 2.42 g Tris, 9 g NaCl, 850-900 µl pure acetic acid into 1000 ml distilled water, adjust pH to 7.2-7.6.
- Wash Buffer
 - Add 0.5 ml of TWEEN 20 into 1000 ml of diluent buffer.

IV. PROTOCOL

Note: Mouse IgM refers to the animal origin of the primary antibody, not the origin of the specimen. This Kit must be used on primary antibodies from mouse.

- 1. Run protein sample and molecular weight standard through polyacrylamide gel electrophoresis (PAGE).
- 2. Transfer the protein sample to a nitrocellulose membrane or PVDF membrane.
- 3. Block membrane:
 - a. Dissolve 2g dry protein powder in 100mL Diluent Buffer.
 - b. Immerse the membrane in blocking buffer and incubate at room temperature for 1.5-2 hours or at 4°C overnight with agitation.
- 4. Wash membrane once for 10 minutes in Wash Buffer.



- 5. Incubate membrane with primary antibody:
 - a. Dilute primary antibody in Diluent Buffer. Incubate membrane with primary antibody solution at room temperature for 2 hours or 4°C overnight with agitation.
 - b. Follow the antibody manufacturer's recommendations for optimized concentration.
- 6. Wash membrane in Wash Buffer with gentle agitation, 3 times for 10 minutes each.
- 7. Incubate the membrane with diluted secondary antibody at room temperature for 90 minutes or at 4°C overnight.
 - a. Secondary antibody dilutions typically range from 1:2000-1:10000. Optimal secondary antibody dilutions must be determined empirically.
- 8. Wash membrane in Wash Buffer with gentle agitation, 4 times for 5 minutes each.
- 9. Chemiluminescent Detection:
 - a. Add 50µl chromogenic reagent A and 50µl chromogenic reagent B into 1 ml of distilled water and mix well. Add the working solution onto the membrane and incubate at room temperature until bands appear (usually 30 seconds 5 minutes).
 - b. Note: It is better to prepare the working solution just before use.

V. TROUBLESHOOTING

High Background

Possible Cause	Solution	
Too high antibody concentration	Optimize and decrease antibody concentration	
Aggregate secondary antibody	Filter the secondary antibody through 0.2µm filter	
formation	Use a new secondary antibody	
Too high antibody incubation	Incubate the antibody at 4°C	
temperature		
Non-specific secondary antibody	Run secondary antibody control (without the primary)	
binding or cross-reactivity with	Decrease secondary antibody concentration	
blocking agent		
Cross-reactivity of primary or	Add Tween-20 to the incubation and washing buffer	
secondary antibody with		
blocking agent		
Incompatible blocking agent	Compare different blocking buffers	
Incomplete blocking	Optimize choice of blocking buffer	
	Increase protein concentration in blocking agent	
	Optimize blocking time and/or temperature; Block for 2	
	hours at normal temperature or overnight at 4°C	
	Add 0.05% Tween 20 detergent into blocking agent	
	Add 0.05% Tween 20 detergent into antibody diluents	
	solution	
Insufficient blocking	Extend blocking time or use a compatible blocking agent	
	(e.g. skim milk, BSA, serum, etc.)	
Cross-reactivity of antibody with	Use different blocking agent (Do not use skim milk with	
other proteins	biotin system	
	Reduce secondary antibody concentration	
	Test cross-reactivity between secondary antibody and	
	membrane	



Increase number of washes and buffer volume	
Add 0.05% Tween 20 detergent into washing buffer	
Reduce exposure time	
Use clean tweezers; Operate with gloves	
Use new membranes	
Ensure the liquid is enough to keep the membrane moist	
Use decolorization table in incubation	
Avoid membranes overlapping	
Handle carefully and avoid damaging membrane	
Increase the number of washes	
Nitrocellulose membrane's background is lower than that	
of PVDF membrane	
Make sure membrane is covered with enough liquid and	
prevent it from drying	
uffer Use new buffer or filter buffer before use	
Ensure all equipment and tools are clean and no gel is left	
on membrane	

Weak/No Signal

Possible Cause	Solution	
Improper protein transfer to membrane	Stain gel after transfer is complete to determine transfer is efficient Use Ponceau S to stain membrane to determine transfer is efficient Ensure sufficient contact between gel and membrane during transfer Make sure transfer sandwich is assembled correctly Wet membrane according the instruction Avoid overheating during electro-transfer Use positive control or molecular weight markers Optimize transfer time and current Avoid sample (antigenic determinant) destroy when handling	
Insufficient protein and membrane binding	Adding 20% methanol to transfer buffer Use small-bore membrane	
Insufficient antibody		
Insufficient antigen	Load more protein	
Antigen masking by blocking buffer	Compare different blocking buffers Optimize protein concentration of blocking agent Reduce blocking time	
Presence of sodium azide in buffers	Eliminate sodium azide from buffers	
Too short exposure time	Lengthen film exposure time	
Too short substrate incubation time	Lengthen substrate incubation time to five minutes	
Digestion of protein on membrane	Optimize amount of blocking agent	
Degradation of protein during storage	Re-prepare protein sample	



Make sure primary antibody, secondary antibody, substrate, enzyme system and samples are compatible Use loading control to test effectiveness of second detecting system	
Increase antibody concentration Increase incubation time	
Use mild detergent such as Tween20	
Change blocking agent (commonly used are milk, BSA, serum or gelatin)	
Check instruction Use positive control	
Use positive control Increase loading amount to 20-30 µg protein per well Use protease inhibitor or fractional extract target protein	
Check the transfer with Ponceau S Soak PVDF-membrane in methanol Avoid excessive wash	
Use 0.05% skim milk or no milk diluents buffer Change blocking agent Reduce blocking time	
Prepare fresh antibody and store properly when not in use Avoid repeated freezing and thawing	
Avoid using sodium azide together with HRP- conjugated antibodies	
Mix enzyme conjugate and substrate (no color development when enzyme is inactive) Use activated enzyme conjugate and fresh substrate	
Soak PVDF membrane in 100% methanol	
Use small-bore membrane	
Reduce transfer time	
Try other buffers such as CAPS buffer (pH 10.5) Try low pH value buffers such as acetic acid buffer	
Decrease methanol concentration or use isopropyl alcohol	

RELATED PRODUCTS

Component	Catalog #	Size
10x TBS pH 7.5	MB-012	1000 ml
10x TTBS pH 7.5	MB-013	1000 ml
10X PBS pH 7.2	MB-008	1000 ml
10x PBST pH 7.2	MB-075-1000	1000 ml