



Polink-1 HRP Mouse-NR AEC Detection System for Mouse Primary Antibody

(Polymer-HRP detection system, biotin-free, Anti-mouse primary antibody) Ready-to-use One Step Polymer Detection System Supper clean when using mouse antibody on rat tissue Super Sensitive for AEC Staining

Storage: 4-8°C	Catalog No.	□ D56-110 □ D56-18 □ D56-6	110 ml (bulk, w/o chromogen) 18 ml (with AEC, good for 180 slides) 6 ml (with AEC, good for 50 slides)
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Intended Use:

Detecting MOUSE primary antibody on RAT tissue is a very difficult task in research field due to background issues. Polink-1 HRP Mouse-NR (no-Rat) AEC Detection kit is specially designed to solve the problem. This technology provides excellent specificity to detect mouse primary antibody (user supplied) on rat tissue. Specimen can be frozen or paraffin – embedded tissues, and freshly prepared monolayer cell smears. This detection system is super sensitive when use with AEC chromogen.

Polink-1 HRP Mouse-NR AEC Detection kit is the ONE step polymer detection system that uses polymeric HRP-linked anti mouse secondary antibody to directly detect mouse primary antibody bound to the rat tissue. The secondary antibody was adsorbed to rat, rabbit and human serum proteins. Besides rat tissue Polink-1 HRP Mouse-NR AEC Detection kit also can be used on human tissue and rabbit tissue as well. It is a biotin-free system, therefore, overcomes the non-specific staining caused by streptavidin/biotin system due to endogenous biotin¹. It is a ONE step detection system that is much faster assay compared to traditional two step method (Biotinylated 2nd antibody, and then streptavidin-HRP). These advantages provide laboratories the benefit of more accurate and quicker result, less trouble shooting and better cost-saving.

If user needs most sensitive polymer detection system for mouse primary antibody on rat tissue, one may choose two-step polymer detection system, Polink-2 Plus HRP Mouse-NR AEC kit (Cat No. D59-110, D59-18, D59-6).

Kit components:

Catalog No.	Product Name	Reagent 1: Polymer HRP-linked anti-mouse IgG for AEC (Ready-to-use)	Reagent 2: 2A: substrate buffer (20x) 2B: Chromogen (20x) 2C: H ₂ O ₂ (20x)
D56-110	Polink-1 HRP Mouse-NR Bulk for AEC kit	110ml	Not provided
D56-18	Polink-1 HRP Mouse-NR AEC 18ml kit	18ml	3ml of Reagent 2A 6ml of Reagent 2B 3ml of Reagent 2C
D56-6	Polink-1 HRP Mouse-NR AEC 6ml kit	6ml	2ml of Reagent 2A 4ml of Reagent 2B 2ml of Reagent 2C

Recommended Protocol:

- 1. Fixation: To ensure the quality of the staining and obtain reproducible performance, user needs to supply appropriately fixed tissue and well prepared slides.
- 2. Tissue need to be adhered to the slide tightly to avoid tissue falling off.
- 3. Paraffin embedded section must be deparaffinized with xylene and rehydrated with a graded series of ethanol before staining.
- 4. Cell smear samples should be made as much monolayer as possible to obtain satisfactory results.
- 5. Investigator needs to optimize dilution and incubation times for primary antibodies.
- 6. Three control slides will aid the interpretation of the result: positive tissue control, reagent control (slides treated with Isotype control reagent), and negative control.
- 7. Proceed IHC staining: DO NOT let specimen or tissue dry from this point on.

Staining Procedure	Incubation Time	
	(Min.)	
a. Incubate slides in peroxidese blocking reagent (Ready-to-use 3% H ₂ O ₂ solution) for	10	
10 min.		
b. Rinse the slide using distilled water.		
a. Heat Induced Epitope Retrieval (HIER) may be required for primary antibody	Refer to vendor's data	
suggested by vendor.	sheet	
	 a. Incubate slides in peroxidese blocking reagent (Ready-to-use 3% H₂O₂ solution) for 10 min. b. Rinse the slide using distilled water. a. Heat Induced Epitope Retrieval (HIER) may be required for primary antibody 	

sheet.	b. Wash with PBS 3 times for 2 minutes each time.			
3. Pre-Block (Optional)	a. Add 2 (100 µL) or more drops of 10% Normal Goat Serum to cover the tissue section			
Not provided	and Incubate 10 min.			
-	b. Drain or blot off solution. DO NOT RINSE.			
Primary antibody:	Notes: Investigator needs to optimize dilution and incubation times	30-60		
	a. Apply 2 (100 µL) or more drops of primary antibody to cover the tissue completely.			
Supplied by user	Incubate in moist chamber for 30-60 min.			
	b. Rinse with PBS containing 0.05% Tween-20 3 times for 2 minutes each time.			
Reagent 1: a. Apply 2 (100 µL) or more drops of HRP Polymer-anti-Mouse IgG to cover tissue		10-15		
HRP Polymer-anti-mouse	buse section and Incubate in moist chamber for 10-15 min.			
(x Rat) RTU	c. Rinse with PBS containing 0.05% Tween-20 3 times for 2 minutes each time.			
6. Reagents 2A, 2B and	Reagents 2A, 2B and a. Add 1 drop of Reagent 2A, 1 drop or 2 drop (for high contrast) of Reagent 2B and 1			
2C: AEC Chromogen (20x)	drop of Reagent 2C to 1 mL distilled or deionized water. Mix well. Protect from light and			
	use within one hour.			
	b. Apply 2 drops (100 µL) or enough volume of pre-mixed AEC Chromogen to			
	completely cover tissue. Incubate for 5 min. to 10 min			
	c. Rinse thoroughly with distill water			
Hematoxylin:	a. Counterstain with 2 (100 ul) or more drops hematoxylin to cover tissue completely	20-30 seconds		
	and wait about 20 seconds.			
Supplied by user.	b. Rinse well with tap water for 1-2 min.			
	c. Put slides in PBS until the color turn blue (about 1/2 - 1 min.)			
	d. Rinse in distill water, then rinse well with tap water			
Mounting medium:	Follow the manufacture data sheet procedure for mounting.	Refer to insert		
	Recommended product:			
Supplied by user	1. GB-Mount: Cat. No. E01-18 (18ml), for alcohol soluble substrates (AEC, AP-Red and			
	AP-blue)			
	2. O-Mount: Cat. No. E02-18 (18ml), for DAB and BCIP/NBT			
	3. Simpo-Mount: Cat.No. E03-18 (18ml), or E03-100 (100ml), universal permanent			
	mounting medium. Can be used with or without cover slip			

Protocol Notes:

- 1. The fixation, tissue slide thickness, and primary antibody dilution and incubation time affect results significantly. Investigator needs to consider all factors and determine optimal conditions when interpreting the result.
- 2. Tissue staining is dependent upon the proper handling and processing of tissues prior to staining. Improper tissue preparation may lead to false negative results or inconsistent results.
- 3. Do not mix reagents from different lot.
- 4. Do not allow the slides to dry at any time during staining.

Related Products:

Product	Catalog No.	Size	Product		Catalog No.	Size
Polink-1 HRP Mouse Bulk kit for AEC	D15-110	110ml		*Polink-1 HRP Rat-NM 18ml, 6ml	D36-18 / D36-6	18ml / 6ml
				AEC Kit		
Polink-1 HRP Mouse 18ml, 6ml AEC Kit	D15-18 / D15-6	18ml / 6ml		Polink-1 HRP Broad Bulk kit for AEC	D14-110 / D14-60	110ml/
				(for mouse & rabbit antibody)		60ml
Polink-1 HRP Rabbit Bulk kit for AEC	D16-110	110ml		Polink-1 HRP Broad 18ml, 6ml AEC	D14-18 / D14-6	18ml / 6ml
				Kit (for mouse & rabbit antibody)		
Polink-1 HRP Rabbit 18ml, 6ml AEC Kit	D16-18 / D16-6	18ml / 6ml		AEC Kit	C01-12	12ml
Polink-1 HRP Goat Bulk kit for AEC	D34-110	110ml		GB-Mount (Aqueous)	E01-18	18ml
Polink-1 HRP Goat 18ml, 6ml AEC Kit	D34-18 / D34-6	18ml / 6ml		Simpo-Mount (Aqueous)	E03-100 /E03-18	100ml/18ml
*Polink-1 HRP Rat-NM Bulk kit for AEC	D36-110	110ml				

*Polink-1 HRP Rat-NM kit does not cross react with mouse primary antibody

Precautious:

Please wear gloves and take other necessary precautions.

Remarks:

For research use only.

References:

1. <u>Bisgaard K, Pluzed KP</u>. Use of polymer conjugates in immunohitochemistry: A comparative study of a traditional staining method to a staining method utilizing polymer conjugates. <u>Abstract</u> XXI Intl Cong Intl Acad Pathol and 12th World Cong Acad Environ Pathol. Budapest, Hungry, October 20-25, 1996.

2. Shi ZR. Itzkowitz SH, Kim YS. A comparison of three immunoperoxidase techniques for antigen detection in colorectal carcimoma tissues. J Hitochem 36:317-322,