



Polink DS-RR-Hu/Ms A Kit for Immunohistochemistry Staining Polymer-HRP & AP double staining kit to detect two rabbit primary antibodies on human/mouse tissue with DAB (Brown) and AP-Red+ (Red)

Storage: 2-8°C	Catalog No.: ☐ DS204A-6/D80-6A 12ml*	60 slides**	
	☐ DS204A-18 36ml*	180 slides**	
	☐ DS204A-60 120ml*	600slides**	
	*Total volume of polymer Conjugates		
	** if use 100µl per slide	** if use 100µl per slide	

Intended Use:

Polink DS-RR-Hu/Ms A Kit is designed to use with user supplied two rabbit antibodies to detect two distinct antigens on human tissue or cell samples. This kit has been tested in paraffin tissue. However, this kit can be used on frozen specimen and freshly prepared monolayer cell smears.

Double staining is one of most common methods used in immunohistostaining that allow revealing two distinct antigens in a single tissue^{1, 2}. **Polink DS-RR-Hu/Ms A Kit** from GBI Labs (Golden Bridge International) supplies two polymer enzyme conjugates: HRP polymer anti-Rabbit IgG and AP polymer anti-Rabbit IgG with two distinct substrates/chromogens, DAB (brown color, use with HRP polymer anti-Rabbit IgG) and AP-Red+ (red color, use with AP polymer anti-Rabbit IgG). **Polink DS-RR-Hu/Ms A Kit** is non-biotin system that avoids endogenous biotin non-specific binding.

Kit Components:

Component No.	Content	12ml Kit	36ml Kit	120ml Kit
Reagent 1	HRP polymer anti-Rabbit IgG(RTU)	6 ml	18 ml	60 ml
Reagent 2A	DAB substrate buffer(RTU)	6 ml	18 ml	60 ml
Reagent 2B	DAB chromogen(20X)	1ml	1ml	6ml
Reagent 3A	DS-RR Blocker A	6 ml	18 ml	60 ml
Reagent 3B	DS-RR Blocker B	6 ml	18 ml	60 ml
Reagent 4	AP polymer anti-Rabbit IgG(RTU)	6 ml	18 ml	60 ml
Reagent 5A	AP-Red Plus Enhancer (40x)	1ml	1ml	3ml
Reagent 5B	AP-Red Plus Solution (40x)	1ml	1ml	3ml
Reagent 5C	AP-Red Plus Substrate (20x)	4ml	4ml	12ml
Reagent 6	Simpo-Mount solution(RTU)	6 ml	18 ml	60 ml

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 deparffinized 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. Three control slides will aid the interpretation of the result: positive tissue control, reagent control (slides treated with Isotype control reagent), and negative control.
- 6. It takes about 30 minutes to dissolve AP-Red+ tablet into the substrate buffer. Make sure to start preparing AP-Red+ solution near the end of the secondary antibody incubation.
- 7. Proceed IHC staining: DO NOT let specimen or tissue dry from this point on.

Reagent	Stainin	g Procedure	Incubation Time
			(Min.)
Peroxidase Blocking	a.	Incubate slides in peroxidese blocking reagent (Ready-to-use	10 min
Reagent		3% H ₂ O ₂ solution) for 10 minutes.	
Not provided	b.	Rinse the slide using distilled water.	
2. HIER Pretreatment:	a.	Heat Induced Epitope Retrieval (HIER) may be required for	
Refer to antibody data		primary antibody suggested by vendor.	
sheet.	b.	Wash with PBS containing 0.05% Tween-20 for 2 min.,3 times.	

3. Preblock (optional)	For paraffin section, Improved formula saves the need for a preblock step.	
(For frozen tissue, preblock may or may not be required depending on fixative. (Preblock catalogue No.:E07 was Recommended.)	
4. Rabbit Antibody 1:	Notes: Investigator needs to optimize dilution and incubation times	30-60 min
Supplied by user	prior to double staining.	
	a. Apply 2 drops or enough volume of rabbit primary antibody 1	
	to cover the tissue completely. Incubate in moist chamber for 30-60 min.	
	b. Rinse with PBS containing 0.05% Tween-20 for 2 min., 3	
	times.	
5.Reagent 1:	a. Apply 1drop (50µl) of Reagent 1 HRP polymer anti-Rabbit IgG	20-30 min
HRP polymer anti-Rabbit	to cover each section.	
IgG(RTU)	b. Incubate in moist chamber for 20-30 min.	
	c. Rinse with PBS containing 0.05% Tween-20 for 2 min., 3	
6. Reagents 2A, 2B:	times. a. Add 1 drop or 2 drops (for higher sensitivity and contrast) of	3-10 min
o. Reagents ZA, ZB.	Reagent 2B to 1 ml Reagent 2A . Mix well. Protect from light	3-10 mm
2A: DAB Substrate (RTU)	and use within 5 hours.	
2B: DAB Chromogen(20x)	b. Apply 2 drops or enough volume of DAB CHROMOGEN to	
	completely cover tissue. Incubate for 3-10 min.	
	c. Rinse thoroughly with distilled water 4 times, 2 minutes each	
	time. d. Rinse with PBS containing 0.05% Tween-20 for 2 min., 3	
	times.	
7. Reagent 3A:	a. Apply 2 drops or enough volume of Reagent 3A DS-RR	30 min
DS-RR Blocker A	Blocker A to cover the tissue completely. Mix well on the slide	
	and Incubate in moist chamber for 30 min.	
	b. Rinse with PBS containing 0.05% Tween-20 for 2 min., 3	
8. Reagent 3B:	times. a. Apply 2 drops or enough volume of Reagent 3B DS-RR	5 min
DS-RR Blocker B	Blocker B to cover the tissue completely. Mix well on the slide	3 111111
	and Incubate in moist chamber for 5 min.	
	b. Rinse with PBS containing 0.05% Tween-20 for 2 min., 3	
O. Dahhit antihadı. 2	times. Notes: Investigator needs to optimize dilution and incubation times	20.00
Rabbit antibody 2: Supplied by user	prior to double staining.	30-60 min
	a. Apply 2 drops or enough volume of rabbit primary antibody 2	
	to cover the tissue completely.	
	b. Rinse with PBS containing 0.05% Tween-20 for 2 min., 3	
10 Persont 4:	times.	1E main
10. Reagent 4: AP polymer anti-Rabbit	a. Apply 1drop (50µl) of Reagent 4 AP polymer anti-Rabbit IgG to cover each section.	15 min
IgG(RTU)	b. Incubate in moist chamber for 30 min.	
J = (· · · = /	c. Rinse with PBS containing 0.05% Tween-20 for 2 min., 3	
	times.	
44 B	d. Rinse with tap water.	
11. Reagent 5A, 5B, 5C:	a. Add 1 drop (50µl) of reagent 5A and 1 drop of reagent 5B to a	15-20 min
5A: AP-Red Plus Enhancer (40x)	test tube. Mix well and set at room temperature for 5 minutes. b. Add 2ml of distilled water to the mixture. Mix well.	
5B: AP-Red Plus Solution (40x)	c. Add 4 drops (200µl) of Reagent 5C and mix well.	
5C: AP-Red Plus Substrate	d. Apply 2 drops (100µl) or enough volume of AP-Red Plus	
(20x)	solution to completely cover the tissue. Incubate for 15-20	
	min., observe appropriate color development	
	e. Rinse well with distilled water. (AP-Red Plus is alcohol soluble; do not dehydrate.)	
12. HEMATOXYLIN	a. Counterstain with 2 drops (100µl) or enough volume of	
Not provided	hematoxylin to completely cover tissue. Incubate for 10-15	
_	seconds.	
	b. Rinse thoroughly with tap water for 2-3 min	
	c. Put slides in PBS until show blue color (about ½ - 1 min.) d. Rinse well in distilled water	
	d. Rinse well in distilled water	

13. Reagent 6:	a.	Apply 2 drops (100 µl) or enough volume of Reagent 6 to	30 min. in 40-
Simpo-Mount(RTU)		cover tissue when tissue is wet. Rotate the slides to allow	50°C oven
		Simpo-Mount spread evenly. DO NOT coverslip.	Or:
	b.	Place slides horizontally in an oven at 40-50°C for at least 30	overnight at room
		minutes or leave it at room temperature until slides are	temperature
		thoroughly dried. Hardened Simpo-Mount forms an	
		impervious polymer barrier to organic solvent. Do not use oil	
		directly on the top of dried Simpo-Mount.	

Protocol Notes:

- 1. The fixation, tissue slide thickness, antigen retrieval and primary antibody dilution and incubation time effect results significantly. Investigator needs to consider all factors and determine optimal conditions when interpret the result.
- 2. Simpo-Mount is water-based mounting medium for immunohistology. It may be used as the permanent mounting media. It does not need coverslip. However, if you need to coverslip, after Simpo-Mount dried, dip the slide in xylene and take out immediately. Apply O-Mount (organic mounting solution, Catalog No. E02-15) on the tissue and place cover glass on the slide. Store it after dry completely.

Precautious:

DAB may be carcinogenic. Please wear gloves and take other necessary precautions.

Remarks:

For research use only.

References:

- 1. <u>De Pasquale A, Paterlini P, Quaglino D</u>. *Immunochemical demonstration of different antigens in single cells in paraffinembedded histological sections*. <u>Olin Lab Haematol</u>. 1982;4(3):267-72.
- 2. Polak J. M and Van Noorden S. <u>Introduction to Immunocytochemistry Second Edition</u>. Bios Scientific Publishers. P41-54. 1997