

PolyStain DS Kit - for 2 Mouse antibody on Human tissue

For co-localization (Emerald/Permanent Red)

NB-23-00099- 3(120 ml)

NB-23-00099- 2(36 ml)

NB-23-00099- 1(12 ml)



PolyStain DS Kit - for 2 Mouse antibody on Human tissue For co-localization (Emerald/Permanent Red)

NB-23-00099-1; NB-23-00099-2; NB-23-00099-3

INTENDED USE:

Storage: 2-8ºC

PolyStain DS Kit is designed to use with user supplied two mouse primary 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 the most common methods used in immunohistochemistry to evaluate two distinct antigens in a single tissue. PolyStain DS Kit from NeoBiotech Labs supplies two polymer enzyme conjugates: HRP polymer anti-Mouse IgG and AP polymer anti-Mouse IgG with two distinct substrates/chromogens, Emerald (Green color, use with HRP polymer anti-Mouse IgG) and Permanent-Red (Red color, use with AP polymer anti-Mouse IgG). Simplified steps offer a much faster protocol and the blocking buffers prevent false positives when using two primary antibodies from the same host species. Another advantage of NeoBiotech Kit, it allows the researcher to visualize when two proteins are co-localized because of the color change when the chromogens overlap that can be semi-quantitative. For example, if the area of co-localization stains blue, the antigen indicated by Emerald is at higher concentration in the cell and if the color is purple than the antigen indicated by Permanent-Red is expressed at higher concentrations. PolyStain DS Kit is non-biotin system that avoids endogenous biotin nonspecific binding.

KIT COMPONENTS:

Component No.	Content	12mL Kit	36mL Kit	120mL Kit
Reagent 1	AP Polymer anti-Mouse IgG (RTU)	6ml	18ml	60ml
Reagent 2A	Permanent Red Substrate (RTU)	7ml	18ml	60ml
Reagent 2B	Permanent Red Activator (5x)	1.4ml	3.6ml	12ml
Reagent 2C	Permanent Red Chromogen (100x)		180µL	0.6ml
Reagent 3	eagent 3 Antibody Blocker (40x)		50mL	125mL
Reagent 4A DS-MM Blocker A (RTU)		6mL	18mL	60mL
Reagent 4B DS-MM Blocker B (RTU)		6mL	18mL	60mL
Reagent 5	HRP Polymer anti-Mouse IgG (RTU)		18mL	60mL
Reagent 6	gent 6 Emerald Chromogen (RTU)		18mL	70mL
Reagent 7	U-Mount (RTU)	3mL	9mL	NA



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. Proceed IHC staining: **DO NOT** let specimen or tissue dry from this point on.

Reagent	Staining Procedure	Incubation Time (Min.)
1. Peroxidase and alkaline phosphatase	a. Incubate slides in peroxidase and alkaline phosphatase blocking reagent. We recommend NeoPure Dual Enzyme Block NB-23-	10 min.
Blocking Reagent Not provided	00193 for 10 minutesb. Rinse the slide using distilled water.	
2. HIER Pretreatment: Refer to antibody data sheet.	a. Heat Induced Epitope Retrieval (HIER) may be required for primary antibody suggested by vendor.	
3. Preblock (optional)	b. Wash with PBS containing 0.05% Tween-20 for 2 min., 3 times. For paraffin section, Improved polymer formula saves the need for a pre-block step. However some primary antibodies may still require it, this information should be determined by the user prior to using DS kit. 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. NB-23-00169 was Recommended.)	
4. Mouse Antibody 1: Supplied by user	 Notes: Investigator needs to optimize dilution and incubation times prior to double staining. a. Apply 2 drops or enough volume of mouse 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 	30 - 60 min.



5.Reagent 1	a. Apply 2 drops (100µL) of Reagent 1 (AP polymer anti-Mouse	15 min		
AP polymer anti-Mouse	IgG) to cover each section.			
IgG (RTU)	b. Incubate in moist chamber for 15 min.			
	c. Rinse with PBS containing 0.05% Tween-20 for 2 min., 3 times.			
	d. Rinse well with distilled or tap water.			
6. Reagent 2A, 2B, 2C	a. Add 200µL of Reagent 2B (Activator) into 1mL of Reagent 2A	10 min		
Reagent 2A:	(Substrate buffer) and mix well. Add 10µL of Reagent 6C			
Permanent Red	(Chromogen) into the mixture and mix well. [Note: For fewer			
Substrate (RTU)	slides, Add 100μL of Reagent 2B (Activator) into 500μL of			
Reagent 2B:	Reagent 2A (Substrate buffer) and mix well. Add 5μL of			
Permanent Red	Reagent 2C (Chromogen) into the mixture and mix well.]			
Activator (5x)	b. Apply 2 drops (100µL) or enough volume of Permanent Red			
Reagent 2C:	working solution to completely cover the tissue. Incubate for 10			
Permanent Red	min, observe appropriate color development.			
Chromogen (100x)	c. Rinse well with distilled water.			
7. Reagent 3: Antibody	<u>Note</u> : This step will block antibodies of previous step so no cross	10 min		
Blocker(40x)	reaction will occur at end of protocol. HIER can be done			
(Optional)	immediately after Antibody Blocker step if only one primary			
	antibody requires antigen retrieval. For frozen tissue a lower			
Must test if	temperature of 65°C must be used for Antibody Blocker (Reagent			
antibody/antigen	3) to prevent tissue from dissociating from slide.			
interaction is heat	a. Use hot plate or water bath to heat diluted Reagent 3 to 1x			
sensitive.	solution (1 part of Antibody Blocker in 39 parts of distilled			
	water) to 80°C. Make enough volume to cover the tissue in beaker.			
Please skip this step if	b. Put slides in heated Antibody Blocker for 10 minutes at 80°C.			
antigen retrieval is	c. Remove slides from the Antibody blocker; cool slides 5 seconds.			
used for 2 nd Ms	d. Rinse slides in multiple changes of distilled water. e. Wash with			
Primary Antibody.	PBS/ 0.05% Tween20 for 2 minutes, 3 times.			
8. Reagent 4A	a. Apply 2 drops or enough volume of Reagent 4A (DSMM	30 min		
DS-MM Blocker A	Blocker A) to cover the tissue completely. Mix well on the slide	30 mm		
DO WINI DIOCKCI II	and Incubate in moist chamber for 30 min.			
	b. Rinse with PBS containing 0.05% Tween-20 for 2 min., 3 times			
9. Reagent 4B	a. Apply 2 drops or enough volume of Reagent 4B (DSMM	5 min.		
DS-MM Blocker B	Blocker B) to cover the tissue completely. Mix well on the slide	U		
	and Incubate in moist chamber for 10 min.			
	b. Rinse with PBS containing 0.05% Tween-20 for 2 min., 3 times.			
10. Mouse antibody 2:	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 mouse primary antibody			
	2 to cover the tissue completely.			
	b. Rinse with PBS containing 0.05% Tween-20 for 2 min., 3			
	times			



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11. Reagent 5	a. Apply 2 drops (100μL) of Reagent 5 AP polymer AntiMouse	15 min			
HRP polymer anti-	IgG to cover each section.				
Mouse IgG (RTU)	b. Incubate in moist chamber for 15 min.				
	c. Rinse with PBS containing 0.05% Tween-20 for 2 min, 3 times.				
	d. Rinse with tap water.				
12. Counterstain	a. Dip the slide in diluted hematoxylin for 5 seconds. (You may	5 seconds			
(Optional)	dilute hematoxylin 1:5 in dH_2O). DO NOT over stain with				
	hematoxylin.				
Not provided	b. Rinse thoroughly with tap water for 2min.				
	c. Put slides in PBS for 5 seconds to blue, DO NOT over blue.				
	d. Rinse well in distilled or tap water for 2min.				
	e. Wash with PBS/0.05% Tween20 for 2 min, 3 times.				
13. Reagent 6	a. Apply 1 to 2 drops (50-100μL) of Reagent 6 (Emerald	5 min			
Emerald	Chromogen) to cover the tissue completely.				
Chromogen(RTU)	b. Incubate in moist chamber for 5 minutes.				
	c. Wash slides in tap water for 1 minute.				
	d. Rinse with distilled water.				
	Important to READ: Emerald Chromogen is water soluble, do				
	counter stain first. Do not leave slides sitting in water. Always stain				
	Emerald chromogen AFTER Permanent Red stain because GBI-				
	Permanent Red removes the Emerald and after hematoxylin.				
14.Dehydrate section	Note: Please wipe off extra water and air dry slides before				
	dehydration and clear.				
	a. Dehydrate with 85% ethanol 20seconds.				
	b. Dehydrate with 95% ethanol 20seconds.				
	c. Dehydrate with 100% ethanol 20seconds.				
	d. Dehydrate with 100% ethanol 20seconds.				
	e. Dehydrate with 100% ethanol 20seconds.				
	f. Dehydrate with xylene 20seconds.				
	CAUTION: DO NOT dehydrate with xylene longer than 20				
	seconds! It will erase Permanent Red stain!				
15. Reagent 7 U-	a. Apply 1 to 2 drops (50-100µL) of Reagent 7 (U Mount) to cover				
15. Reagent 7 U- Mount(RTU)					
	a. Apply 1 to 2 drops (50-100µL) of Reagent 7 (U Mount) to cover				
	a. Apply 1 to 2 drops (50-100µL) of Reagent 7 (U Mount) to cover the tissue section and apply glass coverslip.				



TROUBLE SHOOT:

PROBLEM	TIPS			
Uneven stain on 2 primary antibodies	1. Need to adjust the titer of each antibody.			
	2. The amount of each protein expressed on tissue			
	may be different.			
	3. Set slides in water too long so that Emerald is			
	washed away.			
	4. Set slides in Xylene too long so that Permanent			
	Red is washed away			
Emerald Chromogen is blue not green when non	Emerald should be green when not co-localized			
co-localized with Permanent Red.	with Permanent Red. If Emerald chromogen is blue			
	the titer on the primary antibody is not dilute			
	enough for the protocol. Re-titer primary antibodies			
	individually first.			
No stain on 1 or 2 antibodies	Missing steps or step reversed.			
Green Background on the slide	1. Titer primary antibody.			
	2. Use 10% Donkey serum, goat or horse serum as			
	a preblock.			
Permanent Red is leaching	1. Use fresh 100% ethanol and xylene.			
	2. Slide sat too long in xylene. Do not go over			
	20seconds!			
Artifacts on slides	Slides not completely dried before mount. Use fresh			
	100% Ethanol and xylene.			

PRECAUTIONS:

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

FOR RESEARCH USE ONLY



Work Sheet for NB-23-00099 Kit

We designed this work sheet to help you track of each step. We recommend you use this sheet to record the actual time of each step conducted as it will be helpful for questions with our technical support. To insure that all steps are done properly, we recommend that the user fill in the actual time of their experimental step and any variation. Results will vary if time recommendations are not followed.

RTU translates to ready to use.

- Used for tester to check "√" each step during the experiment
- Steps follow after de-paraffinization
- Refer to insert for details of each step

<u>NB-23-00099</u> Protocol is suitable when both mouse and rabbit primary antibodies need or do not need pretreatment step

Protocol Step	NB-23-00097 Protocol	Experiment 1 Date:	Experiment 2 Date:	Experiment 3 Date:	Experiment 4 Date:
Step 1	Peroxidase & Alkaline Phosphatase Block User supplied				
Step 2 (Optional)	HIER if needed User supplied (up to 60 min)				
Step 3 Optional	Preblock if needed User supplied				
Step 4	Mouse 1°Ab #1 User supplied (30-60 min.)				
Step 5	Reagent 1 AP Polymer anti-Mouse IgG RTU(15min) Rinse with distilled water				
Step 6	Reagent 2A,Reagent 2B& Reagent 2C Permanent Red requires mixing (10min)				



Step 7	Reagent 3 Antibody Blocker (10 min)		
Step 8	Reagent 4A DS-MM Blocker A RTU (30 min)		
Step 9	Reagent 4B DS-MM Blocker B RTU (5 min)		
Step 10	Ms 1°Ab #2 User supplied (30-60 min)		
Step 11	Reagent 5 HRP Polymer anti-Mouse IgG RTU (15min) Rinse with tap water		
Step 12	Counter stain (Do not over counter stain) Hematoxylin User supply Wash with PBS/0.05% Tween20 for 2 min, 3 times		
Step 13	Reagent 6 Emerald Chromogen RTU (5min)		
Step 14	Dehydrate section 20seconds for each step It is important to follow the protocol.		
Step 15	Reagent 7 U-Mount RTU Mount & coverslip		
Result	Stain pattern on controls are correct: Fill in Yes or NO		