

PolyStain DS Kit - for 2 Mouse antibody on Rodent tissue

(DAB/Permanent Blue)

NB-23-00101- 3(120 ml) NB-23-00101- 2(36 ml) NB-23-00101- 1(12 ml)





PolyStain DS Kit - for 2 Mouse antibody on Rodent tissue (DAB/Permanent Blue)

NB-23-00101-1; NB-23-00101-2; NB-23-00101-3

Storage: 2-8ºC

INTENDED USE:

The PolyStain DS Kit is designed to use with two user supplied mouse antibodies to detect two distinct antigens on mouse and rat tissue or cell samples. Specimens can be frozen or paraffin embedded, or freshly prepared monolayer cell smears. Double staining is a common method used in immunohistochemistry that allows for detection of two distinct antigens in a single tissue. This kit uses an HRP or AP polymer based technology combined with a proprietary blocking buffer system that achieves ultra-sensitivity with no background or cross reactivity. PolyStain DS Kit from NeoBiotech labs supplies the user with primer system to enhance the two polymer enzyme conjugates anti-mouse IgG HRP-polymer and anti-mouse IgG AP-polymer with two distinct substrates/chromogens, Permanent Red and DAB. Permanent Red reacts with anti-mouse IgG HRP-polymer conjugate to produce a red color. DAB chromogen reacts with anti-mouse IgG HRP-polymer conjugate to produce a brown color. PolyStain DS Kit is a non-biotin system that avoids the extra steps involved in blocking non-specific binding due to endogenous biotin. Please read the protocol carefully and use the experimental record sheet to keep track of your progress throughout the protocol.

Component No.	Content	12mL Kit	36mL Kit	120mL Kit
Reagent 1	Mouse Primer (RTU)	6mL	18mL	60mL
Reagent 2	Mouse HRP Polymer (RTU)	6mL	18mL	60mL
Reagent 3A	DAB Substrate (RTU)	12mL	15mL x 2	70mL
Reagent 3B	DAB Chromogen (20x)	1.5mL	2mL	3.5mL
Reagent 4	Antibody Blocker (40x)	2 x 15mL	50mL	100mL
Reagent 5A	DS-MM Blocker A (RTU)	6mL	18mL	60mL
Reagent 5B	DS-MM Blocker B (RTU)	6mL	18mL	60mL
Reagent 6	Mouse AP Polymer (RTU)	6mL	18mL	60mL
Reagent 7A	Permanent Red Substrate (RTU)	15mL	18mL x 2	70mL
Reagent 7B	Permanent Red Activator (5x)	3mL	7.2mL	14mL
Reagent 7C	Permanent Red Chromogen (100x)	150µL	360µL	0.7mL
Reagent 8	NeoMount Universal	6mL	18mL	70mL

KIT COMPONENTS:



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 with IHC staining: **DO NOT** let specimen or tissue dry from this point on.
- 7. <u>Note</u>: We recommend TBS-T to be used as the wash buffer to get the highest sensitivity and clean background. Phosphate in the PBS-T may inhibitor the activity of the alkaline phosphatase.

Reagent	Staining Procedure	Incubation Time (Min.)
1. Peroxidase and alkaline phosphatase Blocking Reagent Supplied by user	 a. Incubate slides in peroxidase and alkaline phosphatase blocking reagent (NeoPure Dual Enzyme Block NB-23-00193 is Recommended) for 10 minutes. b. Rinse the slides using 2 changes of distilled water. 	10 min.
2. HIER Pretreatment: Refer to antibody data sheet.	 a. Heat Induced Epitope Retrieval (HIER) may be required for primary antibody. Refer to antibody datasheet. b. Wash with PBS-T containing 0.05% Tween-20 or 1X TBS-T (See note 8 above); 3 times for 2 minutes each. 	60 – 90 min.
3. Ms Primary Antibody1:Supplied by user	 <u>Notes</u>: Investigator needs to optimize dilution and incubation times prior to double staining. Should use as dilute as possible to prevent cross reaction. 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. Wash with PBS-T containing 0.05% Tween-20 or 1X TBS-T; 3 times for 2 minutes each 	30 – 60 min
4.Reagent 1: Mouse Primer (RTU)	 a. Apply 1-2 drops Reagent 1 (Mouse Primer) or enough to cover each section. b. Incubate in moist chamber for 10 min. c. Wash with PBS-T containing 0.05% Tween-20 or 1X TBS-T; 3 times for 2 minutes each. 	10 min



5.Reagent 2:	a. Apply 1-2 drops Reagent 2 (Mouse HRP Polymer) to cover each section.	10 min
Mouse HRP Polymer	b. Incubate in moist chamber for 10 min.	
(RTU)	c. Wash with PBS-T containing 0.05% Tween-20 or 1X TBS-T ; 3	
(((())))	times for 2 minutes each.	
6. Reagent 3A&	Note: Although the DAB step can be done at the end of protocol,	5 min
Reagent 3B:	we find the DAB chromogen acts as additional shielding between	
C	the first mouse and second mouse. We recommend you do this step	
	now.	
3A: DAB Substrate	a. Add 1 drop of Reagent 3B (DAB Chromogen) to 1mL Reagent	
(RTU)	3A (DAB Substrate). Mix well. Protect from light and use within	
3B: DAB	7 hours.	
Chromogen(20x)	b. Apply 2 drops or enough volume of DAB chromogen mixture to	
	completely cover tissue. Incubate for 5 min.	
	c. Rinse well with distilled water.	
7. Reagent 4:	Note: This step will block antibodies of previous step so no cross	10 min.
Antibody Blocker (40x)	reaction will occur at end of protocol.	
(Optional)	a. Use hot plate or water bath to heat diluted Reagent 4 to 1x	
	solution (1 part of Antibody Blocker in 39 parts of distilled	
Must test if	water) to 80-95°C. Make enough volume to cover the tissue in	
antibody/antigen	beaker.	
interaction is heat	b. For paraffin embedded tissue, put slides in heated Antibody	
sensitive.	Blocker for 10 minutes at 95°-100°C. For frozen embedded	
	tissue, put slides in heated Antibody Blocker for 10 minutes at	
	80°C.	
Please skip this step if	c. Cool slides to 55°C.	
antigen retrieval is used	d. Rinse slides in multiple changes of distilled water.	
for 2 nd Ms Primary		
Antibody after step 8.		20
8. Reagent 5A:	a. a. Wash with PBS-T containing 0.05% Tween-20 or 1X TBS-T ;	30 min.
	3 times for 2 minutes each.	
DS-MM Blocker A	b. Apply 2 drops or enough volume of Reagent 5A (DS-MM	
(RTU)	Blocker A) to cover the tissue completely. Mix well on the slide and Incubate in moist chamber for 30 min.	
	c. Wash with PBS-T containing 0.05% Tween-20 or 1X TBS-T ; 3 times for 2 minutes each.	
11 Descent 5P.		5 000
11. Reagent 5B:	a. a. Apply 2 drops or enough volume of Reagent 5B (DS-MM Blocker B) to cover the tissue completely. Mix well on the slide	5 sec
DS-MM Blocker B	and Incubate in moist chamber for 5 min.	
(RTU)	 b. Wash with PBS-T containing 0.05% Tween-20 or 1X TBS-T; 3 	
$(\mathbf{R}\mathbf{I}\mathbf{U})$	times for 2 minutes each.	
	times for 2 minutes each.	



12. Ms Primary	<u>Notes</u> : Investigator needs to optimize dilution and incubation times	30 - 60 min
Antibody 2:	prior to double staining.	
	a. Apply 2 drops or enough volume of mouse primary antibody 2	
Supplied by user	to cover the tissue completely.	
	b. Wash with PBS-T containing 0.05% Tween-20 or 1X TBS-T ; 3	
	times for 2 minutes each.	
13. Reagent 6:	a. Apply 1-2 drops Reagent 6 (Mouse AP Polymer) or enough to	15 min
	cover each section.	
Mouse AP Polymer	b. Incubate in moist chamber for 15 min.	
	c. Wash with PBS containing 0.05% Tween-20 for 3 times for 2	
	min each. d. Wash with 1X TBS-T only; 3 times for 2 minutes	
	each.	
14. Reagent 7A, 7B, 7C	a. Add 200µL of Reagent 7B (Activator) into 1mL of Reagent 7A	10 min.
	(Substrate) and mix well. Add 10µL of Reagent 7C	
Reagent 7A: Permanent	(Chromogen) into the mixture and mix well. [Note: For fewer	
Red Substrate (RTU)	slides, Add 100µL of Reagent 7B (Activator) into 500µL of	
	Reagent 7A (Substrate) and mix well. Add 5μ L of Reagent 7C	
Reagent 7B: Permanent	(Chromogen) into the mixture and mix well.]	
Red Activator (5x)	b. Apply 2 drops (100μ L) or enough volume of Permanent Red	
	working solution to completely cover the tissue. Incubate for 10	
Reagent 7C: Permanent	min, observe appropriate color development.	
Red Chromogen (100x)	c. Rinse well with distilled water.	
15. HEMATOXYLIN	a. Counterstain with 2 drops $(100\mu L)$ or enough volume of	5 min.
	hematoxylin to completely cover tissue. Incubate for 10-15	
Not provided	seconds.	
	b. Rinse thoroughly with tap water for 2-3 min	
	c. Put slides in PBS or Tris pH 7.4 to 8.4 until blue color appears.	
	d. Rinse well in distilled water	
16. Reagent 8:	a. Apply 2 drops (100μ L) or enough volume of Reagent 8	30 min.
	(NeoMount Universal) to cover tissue when tissue is wet. Rotate	50°C oven
NeoMount Universal	the slides to allow NeoMount Universal spread evenly.	or
(RTU)	b. Place slides horizontally in an oven at 40-50°C for at least 30	overnight at
	minutes or leave it at room temperature until slides are	room
	thoroughly dried.	temperature



PROTOCOL NOTES:

- 1. The fixation, tissue slide thickness, antigen retrieval 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. Permanent Red is insoluble in organic solvent and can be coverslipped as well. However the dehydration steps must be shorter for optimal tissue structure and chromogen signal maintenance.

Note: Please wipe off extra water and air dry slides before dehydration and clear.

- a. 1x 80% Ethanol 20 seconds;
- b. 1x 95% Ethanol 20 seconds;
- c. 3x 100% Ethanol 20 seconds each;
- d. 1x 100% Xylene 20 seconds;
- Add 1 drop of Xylene based mountant (Cat. No. NeoMount Perm NB-23-00156) and coverslip.
 Press to push the air bubble out.

CAUTION: DO NOT dehydrate in Xylene longer than 20 seconds! It will erase Permanent Red stain!

PRECAUTIONS:

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

FOR RESEARCH USE



Work Sheet for NB-23-00101 Kit

We designed work sheet to help you track each step. You may use this sheet for our technical support staff to review if needed. To insure that all steps are done properly, we recommend that the user fill in the actual time of their experimental step. Results will vary if time recommendations are not followed. RTU translates to ready to use.

- Used for tester to check " $\sqrt{}$ "each step during the experiment
- Steps follow after de-paraffinization
- Refer to insert for details of each step

NB-23-00101 Protocol-1 is suitable for:

1) Both mouse primary antibodies need pre-treatment;

2) One mouse primary antibody needs pre-treatment and the other one is not sensitive to pretreatment

Protocol Step	NB-23-00101 Protocol-1 Reagent/Time	Experiment 1 Date:	Experiment 2 Date:	Experiment 3 Date:	Experiment 4 Date:
Step 1	Peroxidase Block User supplied				
Step 2 Optional	HIER if needed User supplied (up to 60 min)				
Step 3	Ms 1°Ab #1 User supplied (30-60 min)				
Step 4	Reagent 1 Ms Primer RTU (10 min)				
Step 5	Reagent 2 Ms HRP Polymer RTU (10 min)				
Step 6	Reagent 3A & 3B DAB Requires mixing! (5 min)				



Step 7	Reagent 4		
	Antibody Blocker(40x) (10 min)		
Step 8	Reagent 5A:		
	DS-MM Blocker A RTU (30 min)		
Step 19	Reagent 5B:		
	DS-MM Blocker B RTU (5 min)		
Step 10	Ms 1°Ab #2		
	User supplied (30-60 min)		
Step 11	Reagent 6 Ms AP Polymer RTU (15 min)		
	Wash with 1xTBS-T only.		
Step 12	Reagent 7A, 7B, & 7C		
	Permanent Red requires mixing (10min)		
Step 13	Counter stain Hematoxylin		
	User supplied		
Step 14	Reagent 8 NeoMount Universal RTU		
	Do not coverslip!		
Result	Stain pattern on controls are correct: Fill in Yes or NO		

Testing result:



NB-23-00101 Protocol-2 is suitable for one mouse primary antibody needs pre-treatment, the other mouse primary antibody is sensitive to pre-treatment.

Protocol Step	NB-23-00101 Protocol-2 Reagent/Time	Experiment 1 Date:	Experiment 2 Date:	Experiment 3 Date:	Experiment 4 Date:
Step 1	Peroxidase Block User supplied				
Step 3	Ms 1°Ab #1 User supplied (30-60 min)				
Step 4	Reagent 1 Ms Primer RTU (10 min)				
Step 5	Reagent 2 Ms HRP Polymer RTU (10 min)				
Step 6	Reagent 3A & 3B DAB Requires mixing! (5 min)				
Step 2	HIER (10-15 min) Cool down (45-60 min) User supplied Skip antibody blocker step 9 if HIER is done since they will achieve same goal.				
Step 8	Reagent 5A: DS-MM Blocker A RTU (30 min)				



Step 9	Reagent 5B:		
	DS-MM Blocker B RTU (5		
	min)		
Step 10	Ms 1°Ab #2		
	User supplied (30-60 min)		
Step 11	<u>Reagent 6</u> Ms AP Polymer RTU (15		
	min)		
	Wash with 1xTBS-T only.		
Step 12	Reagent 7A, 7B, & 7C		
	Permanent Red requires		
	mixing (10min)		
Step 13	Counter stain		
	Hematoxylin		
	User supplied		
Step 14	Poogont 9		
51ep 14	<u>Reagent 8</u> NeoMount Universal RTU		
	Do not covorslin!		
	Do not coverslip!		
Result	Stain pattern on controls are correct: Fill in Yes or		
	NO		

