

NB-23-00090-1

NB-23-00090-2

NB-23-00090-3



NeoStain Poly DS Kit - for Mouse and Rabbit antibody on Human tissue (DAB/Permanent Red)

#Cat: NB-23-00090-1 Size: 6ml #Cat: NB-23-00090-2 Size: 36ml #Cat: NB-23-00090-3 Size: 120ml

Storage: 2-8ºC

Intended use:

The NeoStain Poly DS Kit is designed to use with user supplied mouse and rabbit antibody to detect two distinct antigens on human tissue or cell samples. This kit has been tested in paraffin tissue, but may also be used on frozen specimens and freshly prepared monolayer cell smears.

Double staining is one of most common methods used in immunohistostaining for revealing two distinct antigens in a single tissue 1, 2. NeoStain Poly DS Kit from Golden Bridge International supplies two polymer enzyme conjugates: AP-Polymer anti-Mouse IgG and HRP-Polymer anti-Rabbit IgG with two distinct chromogens, DAB (brown) and Permanent Red(red). User may apply the two enzyme conjugates onto the specimen at the same time and mix them on the slide. Simplified steps allow for a much faster protocol than sequential procedures. NeoStain Poly DS Kit is a non-biotin system that avoids endogenous biotin non-specific binding.

Kit components:

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Component No.	Content	12mL Kit	36mL Kit	120mL Kit	
Reagent 1	HRP Polymer anti-Rabbit IgG (RTU)	6mL	18ml	60ml	
Reagent 2	AP Polymer anti-Mouse IgG(RTU)	6mL	18ml	60ml	
Reagent 3A	DAB Substrate (RTU)	12mL	18mLx2	120ml	
Reagent 3B	DAB Chromogen (20x)	1.5mL	2mL	6ml	
Reagent 4A	Permanent Red Substrate (RTU)	15mL	18mLx2	12mLx2	
Reagent 4B	Permanent Red Activator (5x)	3mL	7.2mL	12mLx2	
Reagent 4C	Permanent Red Chromogen (100x)	150μL	360μL	1.2mL	
Reagent 5	Simpo-Mount	12mL	18mLx2	120mL	

HRP = Horseradish Peroxidase AP = Alkaline Phosphatase Ms = Mouse Rb = Rabbit



Protocol Notes:

- 1. Proper 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 needs to be adhered to the slide tightly to avoid falling off.
- 3. Paraffin embedded sections must be deparaffinize with xylene and rehydrated with a graded series of alcohols before staining.
- 4. Cell smear samples should be prepared as close to a monolayer as possible to obtain satisfactory results.
- 5. Control slides are recommended for interpretation of results: positive, reagent (slides treated with Isotype control reagent), and negative control.
- 6. **DO NOT** let specimen or tissue dry during protocol. This will generate false positive and/or false negative signal.
- 7. The fixation, tissue section thickness, antigen retrieval and primary antibody dilution and incubation time effect results significantly. Investigator needs to consider all factors and determine optimal conditions when interpreting results.
- 8. 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 inhibit the activity of the alkaline phosphatase. Note: 1X TBS-T =50mM Tris HCl, 150mM NaCl, 0.05% Tween-20 pH7.6. Neo Biotech sells 10xTBS-T for your convenience (NB-23-00201)

Reagent	Staining Procedure	Incubation Time (Min.)
1. Peroxidase and Alkaline Phosphatase Blocking Reagent Not Provided We recommend using NeoPure Dual Enzyme Block NB-23-00193 Fast, easy and it will block endogenous alkaline phosphatase	 a. Incubate slides in peroxidase and alkaline phosphatase blocking reagent. We recommend Neopure Dual Enzyme Block NB-23-00193. b. Rinse the slide using distilled water 	10min
2. Antigen retrieval if needed: Refer to primary antibody data sheet.	 a. Refer to primary antibody data sheet for antigen retrieval methods. b. Wash with PBS-T containing 0.05% Tween-20 or 1X TBST (See note 8 above); 3 times for 2 minutes each. 	Up to 1h
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.: NB-23-00169 was Recommended.)	
4 . Primary Antibody Mix: Mix one Mouse and one Rabbit primary antibody Supplied by user.	Note: Investigator needs to optimize dilution prior to triple staining. a. Apply 2 drops or enough volume of mouse and rabbit primary antibody mixture to cover the tissue completely. Incubate in moist chamber for 30-60 min. Recommend 30min to shorten total protocol time. b. Wash with PBS-T containing 0.05% Tween-20 or 1X TBS-T; 3 times for 2 minutes each.	30-60min



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5.Reagent 1 and 2	Note : Make sufficient polymer mixture by adding	
Reagent 1: HRP Polymer anti-Rabbit	Reagent 1 HRP Polymer anti-Rabbit IgG and Reagent 2	
lgG(RTU)	AP Polymer anti-Mouse IgG at 1:1 ratio, mix well. Do Not	30min
Reagent 2: AP Polymer anti Mouse	Mix More than you need for the experiment because	
IgG(RTU)	the polymer mixture may not be as stable as non-mixed	
	polymer.	
	a.Apply 1 to 2 drops (50-100µL) of the mixture to cover	
	the tissue completely.	
	b.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.	
6. Reagent 3A and 3B	Note: Make enough DAB mix by adding 1 drop of	
Reagent 3A:	Reagent 3B (DAB Chromogen) in 1mL of Reagent 3A	
DAB Substrate(RTU)	(DAB Substrate). Mix well. Use within 7 hours.	5min
Reagent 3B:	a. Apply 1 to 2 drops (50-100μL) of your DAB mixture to	
DAB Chromogen(20x)	cover the tissue completely.	
	b. Incubate for 5min.	
	c. Rinse slides in multiple changes of distilled water 3	
	times, 2 each time or under running tap water for 1	
	minute.	
	d. Wash with 1X TBS-T only; 3 times for 2 minutes each.	
7.Reagent 4A, 4B, 4C	Note: Shake Permanent Red Activator before adding	
Reagent 4A:	into Permanent Red Substrate.	
Permanent Red Substrate (RTU)	a. Add 200µL of Reagent 4B (Activator) into 1mL of	
Reagent 4B:	Reagent 4A (Substrate buffer) and mix well. Add 10μL	
Permanent Red Activator (5x)	of Reagent 4C (Chromogen) into the mixture and mix	10 min
Reagent 4C:	well.	
Permanent Red Chromogen (100x)	[Note: For fewer slides, Add 100µL of Reagent 4B	
(To get maximum sensitivity of AP	(Activator) into 500μL of Reagent 4A (Substrate buffer)	
polymer, Please repeat chromogen	and mix well. Add 5μL of Reagent 4C (Chromogen) into	
step)	the mixture and mix well.].	
.,	b. Apply 2 drops (100µL) or enough volume of Permanent Red working solution to completely cover	
	the tissue. Incubate for 10 min, observe appropriate	
	color development. To increase AP signal aspirate or	
	tap off chromogen and apply 2-3 drops (100μL) again	
	of the Permanent Red working solution to completely	
	cover the tissue for additional 5 to 10min.	
	c. Rinse well with distilled water.	
8.HEMATOXYLIN	a. Counterstain with 2 drops (100µl) or enough volume	
Not provided	of hematoxylin to completely cover tissue. Incubate for	
	5 seconds. DO NOT over stain with hematoxylin.	
	b. Rinse thoroughly with tap water for 1 minute.	
	c. Put slides in PBS for 5 seconds to blue, DO NOT over	
	blue.	
	d. Rinse well in distilled or tap water for 1 minute.	



9. Reagent 5 : Simpo-Mount(RTU)	a. Apply 2 drops (100µL) or enough volume of Reagent 5 Simpo-Mount to cover tissue when tissue is wet. Rotate the slides to allow Simpo-Mount spread evenly. b. Place slides horizontally in an oven at 40-50°C for at least 30 minutes or leave it at room temperature until slides are thoroughly dried.	
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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. I-Permanent Red is insoluble in organic solvent and can be coversliped 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;
- e. Add 1 drop of xylene based mountant (Cat. No. NeoBio Mount Perm (Permanent mount for DAB,BCIP/NBT), 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.

Remarks:

For research use only.

References:

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



Work Sheet for NB-23-00090 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 "V" each step during the experiment
- Steps follow after de-paraffinization
- Refer to insert for details of each step

NB-23-00090 Protocol is suitable when both mouse and rabbit primary antibodies need or do not need pre-treatment step

Step /	NB-23-00090 Protocol	Experiment 1	Experiment 2	Experiment 3	Experiment 4
Protocol	Reaction time	Date:	Date:	Date:	Date:
Step 1	Peroxidase or Alkaline				
	Phosphatase Block				
	User supplied				
Step 2	HIER if needed				
(optional)	User supplied (up to 60 min)				
Step 3	Mouse 1°Ab & Rabbit				
	1°Ab mixture				
	(30-60 min.)				
Step 4	Reagent 1& Reagent 2				
	HRP Polymer anti-Rabbit IgG				
	and AP Polymer anti-Mouse				
	IgG require mixing (30min)				
Step 5	Reagent 3A & Reagent 3B				
	DAB Requires mixing!				
	(5 min.)				
Step 6	Reagent 4A, Reagent 4B&				
	Reagent 4C				
	Permanent Red Requires				
	mixing! (10 min)				
Step 7	Counter stain(10-15sec)				
	User supplied				
Step 8	Reagent 5				
	Simpo Mount (RTU)				
Step 11	Dehydrate section				
	20seconds for each step				
	It is important to follow the protocol.				
Result	Stain pattern on controls are				
	correct: Fill in Yes or NO				

The result: