



# NeoSplink HRP Broad Spectrum kit with DAB

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**NB-23-00001-3 (110 ml)**

**NB-23-00001-4 (60 ml)**

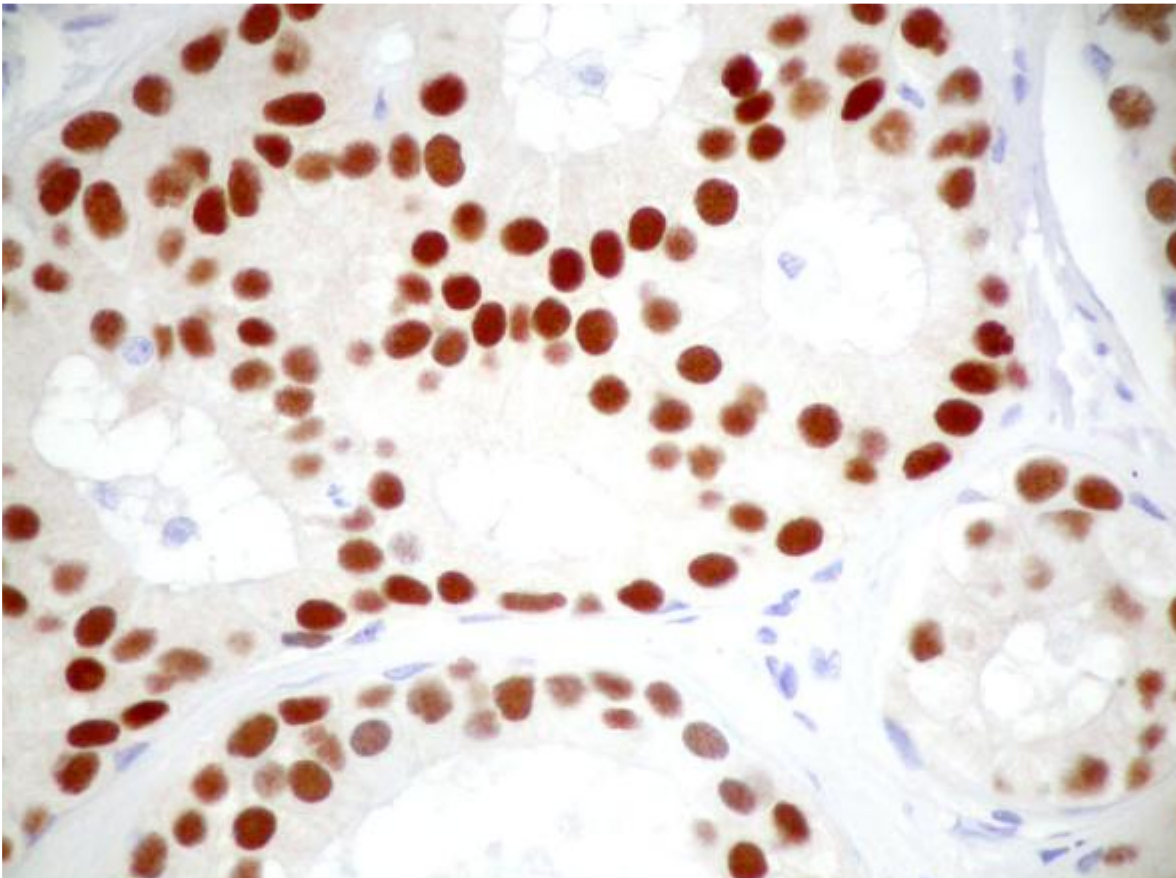
**NB-23-00001-5 (18 ml)**

**NB-23-00001-6 (6 ml)**



## **NeoSplink HRP Broad Spectrum kit with DAB**

*Cat # NB-23-00001-3 ; Cat # NB-23-00001-4; Cat # NB-23-00001-5;  
Cat # NB-23-00001-6*



**Human breast cancer stained by Ms x ER**

### **Intended Use**

NeoSplink HRP Broad Spectrum kit with DAB is intended for using with mouse and rabbit primary antibody (user-supplied) to detect the presence of antigens in human tissue or cell preparations under light microscopy. Most commonly used specimens for this system are: frozen tissue, paraffin-embedded tissue, freshly prepared lymphocytes and fixed culture cells. Horseradish peroxidase (HRP) labeled-streptavidin and biotinylated secondary antibody amplification system has become a standard technique in immunochemical staining<sup>1,2</sup>. NeoSplink HRP Broad Spectrum kit with DAB uses human-adsorbed, biotinylated, affinity-purified secondary antibody reacts with the user supplied primary antibody bound to the specific epitope of the antigen in tissue or cell. Horseradish peroxidase (HRP) labeled streptavidin then reacts with biotinylated secondary antibody to form a HRP-streptavidin-biotin complex.

The HRP enzyme of the streptavidin complex catalyzed the substrate/chomogen, 3,3' diaminobenzidine (DAB substrate) or 3-Amino-9-ethylcarbazole (AEC substrate) reaction to form brown (if use DAB) or red color (if use AEC) deposit at the antigen site. The antigen then can be visualized under microscope. Compared to traditional ABC method which uses avidin, NeoSplink HRP Broad Spectrum kit with DAB demonstrates stronger binding strength to bind biotin and less non-specific background staining.

Higher sensitivity and lower background give NeoSplink HRP Broad Spectrum kit with DAB a higher signal-noise ratio. NeoSplink HRP Broad Spectrum kit with DAB provides users cost effective method for their research. End users may use DAB (Cat. No. NB-23-00141-2) or DAB+ kit (NB-23-00148-1) or AEC (NB-23-00140) chromogen.

### Kit Components

Cat. No.	No. Description	Reagent 1	Reagent 2	Reagent 3	Reagent 4A, B
		Pre-Blocking Solution	Biotinylated second antibody broad spectrum	Streptavidin-peroxidase conjugate	4A: DAB Substrate 4B: DAB Chromogen
NB-23-00001-3	Neo SPlink HRP Broad Bulk Kit	110 ml	110 ml	110 ml	Not included
NB-23-00001-4	Neo SPlink HRP Broad 60ml Kit	60ml	60ml	60ml	Not included
NB-23-00001-5	Neo SPlink HRP Broad DAB 18ml Kit	18ml	18ml	18ml	4A: 15ml x2 4B: 2ml
NB-23-00001-6	Neo SPlink HRP Broad DAB 6ml Kit	6ml	6ml	6ml	4A: 12ml 4B: 1.5ml

### 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 (slide treated with Isotype control reagent), and negative control.
6. Start staining procedures: DO NOT let specimen or tissue dry from this point on.

Reagent	Staining Procedures	Incubation Time (Min.)
1. Peroxidase blocking reagent: Supplied by user.	a. Apply 2 drops (100 $\mu$ L) or enough volume of Peroxidase blocking reagent (Ready-to-use 3% H <sub>2</sub> O <sub>2</sub> solution) to cover the tissue section and incubate b. Rinse the slide using distilled water.	10 min.
2. HIER Pretreatment: refer to antibody spec. sheet	a. Heat Induced Epitope Retrieval (HIER) may be required for primary antibody suggested by vendor b. Wash with PBS 2 min., 3 times.	
3. <b>Reagent 1:</b> Pre-blocking Solution	a. Add 2 drops or enough of volume Pre-blocking Solution to completely cover the tissue section and Incubate b. Blot off solution. DO NOT RINSE.	10 min.
4. Primary antibody: Supplied by user. Investigator needs to optimize dilution and incubation time.	a. Apply 2 drops or enough volume of Primary antibody to cover the tissue section completely. Incubate in moist chamber for 30-60 min. b. Rinse with PBS for 2 min., 3 times.	30-60 min.
5. <b>Reagent 2:</b> Ready to use Secondary antibody	a. Apply 2 drops or enough volume of secondary antibody to cover the tissue section completely and incubate. b. Rinse with PBS for 2 min., 3 times.	10 min.
6. <b>Reagent 3:</b> Ready to use HRP-Streptavidin	a. Apply 2 drops or enough volume of HRP-Streptavidin to cover the tissue section completely and incubate. b. Rinse with PBS for 2 min., 3 times.	10 min.
7. <b>Reagent 4:</b>  <b>4A:</b> DAB Substrate <b>4B:</b> DAB Chromogen concentrate (chromogen may be supplied by user)	a. Adding 1 drop or 2 drops (for higher sensitivity and contrast) of DAB chromogen concentrate ( <b>Reagent 4B</b> ) in 1ml of DAB substrate buffer ( <b>Reagent 4A</b> ). Mix well. b. Apply 2 drops (100 $\mu$ L) or enough volume of pre-mixed DAB Chromogen to completely cover tissue. Incubate for 5 min. Use the prepared DAB solution within 5 hours. c. When appropriate color is developed, rinse under tap water gently for about 1-2 minutes.	5 min.
8. Hematoxylin: Supplied by user	a. Counterstain with 2 drops or enough volume to cover tissue completely and wait about 10-20 seconds. b. Rinse thoroughly under tap water for 1-2 min. c. Put slides in PBS until show blue color (about 30-60 seconds) d. Rinse well in distilled water	
9. Mounting media: Supplied by user	Follow the manufacture data sheet procedure for mounting. Recommended product: 1. Neo-Moun, for alcohol soluble substrates (AEC, AP-Red and AP-blue) 2. Neo O-Mount, for DAB 3. Neo Simpo-Mount, universal permanent mounting medium	



## 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. 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
Neo SPlink HRP Mouse Bulk kit	NB-23-00003-2	110ml	Neo Simplified Streptavidin HRP Rabbit concentrate kit (1:100)	NB-23-00010	1ml
Neo SPlink HRP Mouse DAB Kit	NB-23-00003-3/ NB-23-00003-4	18ml / 6ml	Neo Simplified Streptavidin HRP Mouse concentrate kit (1:100)	NB-23-00011	1ml
Neo SPlink HRP Rabbit Bulk kit	NB-23-00005-2	110ml	Neo Streptavidin Peroxidase (RTU)	NB-23-00026-2/ NB-23-00026-3	110ml 18ml
Neo SPlink HRP Rabbit DAB Kit	NB-23-00005-3/ NB-23-00005-4	18ml / 6ml	Neo SPlink HRP Broad AEC	NB-23-00007-1/ NB-23-00007-2	18ml / 6ml
Neo SPlink HRP Goat Bulk kit	NB-23-00012-1	110ml	Neo SPlink HRP Mouse AEC	NB-23-00008-1/ NB-23-00008-2	18ml / 6ml
Neo SPlink HRP Goat DAB Kit	NB-23-00012-2/ NB-23-00012-3	18ml / 6ml	SPlink HRP Rabbit AEC	NB-23-00009-1/ NB-23-00009-2	18ml / 6ml

## Precautions

Handle all specimens as potential infectious materials, wear gloves and protection cloth.

**THANK YOU FOR USING NEO BIOTECH PRODUCT!**

