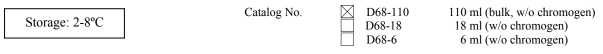




# Polink-2 Plus AP Broad Detection System for Immunohistochemistry

(2-step Polymer-AP detection system, biotin-free, for mouse and rabbit antibody) Polymer Detection System with Super Sensitivity and Specificity



### Intended Use:

Polink-2 Plus AP Broad Detection Kit is the 3rd generation of polymer detection system. It uses mouse and rabbit antibody enhancer to help amplify the polymer-enzyme conjugate reaction to achieve super sensitivity and specificity in immunohistochemistry staining. It produces consistent immunostaining outcomes on archival tissues and on difficult-to-work antibodies. User may need to further dilute primary antibody due to super sensitivity of Polink-2 Plus detection system. It is a biotin-free system, therefore it overcomes the non-specific staining caused by streptavidin/biotin system due to endogenous biotin. Most commonly used specimens for this system are: frozen tissue, paraffin-embedded tissue, freshly prepared lymphocytes and fixed culture cells. It can be used for manual stain or autostainer. Staining conditions need to be optimized by user.

Polink-2 Plus AP Detection System offers a wide choice for primary antibodies, including broad spectrum (for mouse and rabbit primary antibodies), mouse, rabbit, goat, and rat primary antibodies. Refer to **Related Product** section for details.

## Kit components:

Component No.	Content	6mL Kit	18mL Kit	110mL Kit
Reagent 1	Broad Antibody Enhancer(RTU)	6mL	18mL	110mL
Reagent 2	Polymer AP for Mouse & Rabbit(RTU)	6mL	18mL	110mL
Reagent 3A	GBI-Permanent Red Substrate (RTU)	7mL	18mL	NA
Reagent 3B	GBI-Permanent Red Activator (5x)	1.4mL	2x1.8mL	NA
Reagent 3C	GBI-Permanent Red Chromogen (100x)	70µL	180µL	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 deparaffinized with xylene and rehydrated with a graded series of ethanol before staining.
- 4. Cell smear samples should be made into as thin monolayer as possible to obtain satisfactory results.
- 5. Investigator needs to optimize dilution and incubation times for primary antibodies.
- 6. Three control slides will aid the interpretation of the result: positive tissue control, reagent control (slides treated with Isotype control reagent), and negative control.
- 7. Staining steps: DO NOT let specimen or tissue dry from this point on.
- 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. GBI sells 10xTBS-T for your convenience (B11xx)

Reagent	Staining Procedure	Incubation
		Time (Min.)
1. Phosphatase Blocking Reagent Supplied by user		
2. HIER PRETREATMENT:	a. Heat Induced Epitope Retrieval (HIER) may be required for primary antibody suggested by vendor. Please check the data sheet of primary antibody	1hour
	b. Wash with PBS-T containing 0.05% Tween-20 or <b>1X TBS-T(See note 8 above)</b> ; 3 times for 2 minutes each.	
3. PRIMARY ANTIBODY	a. Apply 2 drops ( $100\mu$ L) or enough volume of PRIMARY ANTIBODY to cover the tissue section completely. Incubate in moist chamber for 30-60 min.	30-60min
Supplied by user	b. Wash with PBS-T containing 0.05% Tween-20 or <b>1X TBS-T</b> ; 3 times for 2 minutes each.	
4. Reagent 1 Broad (Mouse/Rabbit)	a. Apply 2 drops $(100\mu L)$ or enough volume of <b>Reagent 1</b> Broad (Mouse/ Rabbit) Antibody Enhancer to cover each section. Incubate in moist chamber for 10 min.	10min
Antibody Enhancer (RTU)	b. Wash with <b>1X TBS-T only</b> ; 3 times for 2 minutes each.	

5. <b>Reagent 2</b> Polymer AP for Mouse/Rabbit (RTU)	<ul> <li>a. Apply 2 drops (100μL) or enough volume of Reagent 2 Polymer AP for Mouse/Rabbit to cover each section. Incubate in moist chamber for 10 min.</li> <li>b. Wash with PBS/Tween(0.05%) 2 min., 3 times or use 1xTBS-T see note #8</li> <li>c. Rinse with tap water.</li> </ul>	10min
6. Reagent3A, 3B, 3C Reagent 3A: GBI-Permanent Red Substrate (RTU) Reagent 3B: GBI-Permanent Red Activator (5x) Reagent 3C: GBI-Permanent Red Chromogen (100x)	<ul> <li>a. Add 200μL of Reagent 3B (Activator) into 1mL of Reagent 3A (Substrate buffer) and mix well. Add 10μL of Reagent 3C(Chromogen) into the mixture and mix well. [Note: For fewer slides, Add 100μL of Reagent 3B (Activator) into 500μL of Reagent 3A (Substrate buffer) and mix well. Add 5μL of Reagent 3C(Chromogen) into the mixture and mix well. ]</li> <li>b. Apply 2 drops (100μL) or enough volume of GBI-Permanent Red working solution to completely cover the tissue. Incubate for 10 min, observe appropriate color development.</li> <li>c. Rinse well with distilled water.</li> </ul>	10 min
<ul> <li>7. Hematoxylin:         <ul> <li>a. Counterstain with 2 (100μL) or more drops hematoxylin to cover tissue compliand wait about 20 seconds.</li> <li>b. Rinse well with tap water for 1-2 min.</li> <li>c. Put slides in PBS until the color turn blue (about ½ - 1 min.)</li> <li>d. Rinse in distill water, then rinse well with tap water</li> </ul> </li> </ul>		20-30 seconds
8. Mounting medium: Supplied by user	<ul> <li>Follow the manufacture data sheet procedure for mounting.</li> <li>Recommended product:</li> <li>1. GB-Mount: Cat. No. E01-18 (18mL), for alcohol soluble substrates (AEC, GBI-Permanent Red and AP-blue)</li> <li>2. Simpo-Mount: Cat.No. E03-18 (18mL), or E03-100 (100mL), universal permanent mounting medium. Can be used with or without cover slip</li> </ul>	Refer to insert

#### **Protocol Notes:**

- 1. The fixation, tissue slide thickness, 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. **GBI-Permanent Red** is insoluble in organic solvent and can be cover sliped 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. O-Mount, E02-18) and coverslip. Press to push the air bubble out.

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

#### **Related Products:**

Product	Catalog No.	Size	Product	Catalog No.	Size
Polink-2 Plus AP Mouse Bulk kit	D69-110	110mL	Polink-2 Plus Mouse-NR AP bulk kit (No cross react to RAT )	D65-110	110mL
Polink-2 Plus AP Mouse 18ml kit / 6ml kit	D69-18 / D68- 6	18mL / 6mL	Polink-2 Plus AP Mouse-NR 18ml/6ml kit (No cross react to RAT)	D65-18 / D65-6	18mL / 6mL
Polink-2 Plus AP Rabbit bulk kit	D70-110	110mL	Fast Red Kit	C03-60	12 Tab + 60mL
Polink-2 Plus AP Rabbit 18ml kit / 6ml kit	D70-18 / D70- 6	18mL / 6mL	Klear Dual Enzyme Block	E36-100 E36-18	100mL 18mL
Polink-2 Plus AP Goat bulk Kit	D66-110	110mL	BCIP/NBT Kit	C05-100 C05-18	100mL/ 18mL
Polink-2 Plus AP Goat 18ml kit / 6ml kit	D66-18 / D66- 6	18mL/ 6mL	GB-Mount (Aqueous)	E01-18	18mL
Polink-2 Plus AP Rat-NM bulk	D67-110	110mL	Simpo-Mount (Aqueous)	E03-100 /E03-18	100mL/ 18mL
Polink-2 Plus AP Rat-NM 18ml kit / 6ml kit	D67-18 / D67- 6	18mL/ 6mL	TBS-T (10X Concentrate)	B11-500 B11-1L	500mL 1000mL

Precautious: Please wear gloves and take other necessary precautions.

**Remarks:** For research use only.