



# Polink-2 Plus AP Rabbit Detection System for Immunohistochemistry

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

Storage: 2-8°C

Catalog No. D70-110 D70-18 D70-6

110 mL (Bulk, w/o chromogen) 18 mL (w/ GBI-Permanent Red) 6 mL (w/ GBI-Permanent Red)

#### Intended Use:

**Polink-2 Plus AP Rabbit Detection Kit** is the 3rd generation of polymer detection system. It uses rabbit antibody enhancer to help amplify the polymerenzyme 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	Rabbit Antibody Enhancer(RTU)	6mL	18mL	110mL
Reagent 2	Polymer AP anti-Rabbit(RTU)	6mL	18mL	110mL
Reagent 3A	GBI-Permanent Red Substrate (RTU)	7mL	18mL	NA
Reagent 3B	GBI-Permanent Red Activator (5x)	1.4mL	3.6mL	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
1. HIER PRETREATMENT:	<ul> <li>a. Heat Induced Epitope Retrieval (HIER) may be required for primary antibody suggested by vendor. Please check the data sheet of primary antibody</li> <li>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.</li> </ul>	
2. PRIMARY ANTIBODY Supplied by user	<ul> <li>a. Apply 2 drops (100 μL) or enough volume of PRIMARY ANTIBODY to cover the tissue section completely. Incubate in moist chamber for 30-60 min.</li> <li>b. Wash with PBS-T containing 0.05% Tween-20 or <b>1X TBS-T</b>; 3 times for 2 minutes each.</li> </ul>	
3. Reagent 1 Rabbit Antibody Enhancer (RTU).	<ul> <li>a. Apply 2 drops (100 μL) or enough volume of <b>Reagent 1</b> Rabbit Antibody Enhancer to cover each section. Incubate in moist chamber for 10 min.</li> <li>b. Wash with PBS/Tween(0.05%) 2 min., 3 times.</li> </ul>	

4. Reagent 2	a. Apply 2 drops (100 $\mu$ L) or enough volume of <b>Reagent 2</b> Polymer AP anti-Rabbit to			
Polymer AP anti-Rabbit (RTU)	cover each section. Incubate in moist chamber for 10 min.			
	b. Wash with <b>1X TBS-T only</b> ; 3 times for 2 minutes each.			
5. Reagent3A, 3B, 3C	a. Add 200µL of <b>Reagent 3B</b> (Activator) into 1mL of <b>Reagent 3A</b> (Substrate buffer) and			
Reagent 3A:	mix well. Add $10\mu L$ of <b>Reagent 3C</b> (Chromogen) into the mixture and mix well.			
GBI-Permanent Red Substrate (RTU) Reagent 3B:	[Note: For fewer slides, Add 100µL of Reagent 3B (Activator) into 500µL of Reagent			
GBI-Permanent Red Activator (5x) Reagent 3C:	<b>3A</b> (Substrate buffer) and mix well. Add 5µL of <b>Reagent 3C</b> (Chromogen) into the mixture and mix well. ]	10min		
GBI-Permanent Red Chromogen (100x)	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.			
	c. Rinse well with distilled water.			
6. Hematoxylin:	a. Counterstain with 2 (100 $\mu$ L) or more drops hematoxylin to cover tissue completely and			
Supplied by user.	wait about 20 seconds.			
	b. Rinse well with tap water for 1-2 min.			
	c. Put slides in PBS until the color turn blue (about $\frac{1}{2}$ - 1 min.)			
	d. Rinse in distill water, then rinse well with tap water			
7. Mounting medium:	Follow the manufacture data sheet procedure for mounting.			
Supplied by user	Recommended product:			
-	1. GB-Mount: Cat. No. E01-18 (18ml), for alcohol soluble substrates (AEC, AP-Red and			
	AP-blue)			
	2. Simpo-Mount: Cat.No. E03-18 (18ml), or E03-100 (100ml), universal permanent			
	mounting medium. Can be used with or without cover slip			

#### **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 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. 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 Mouse-NR AP bulk kit		
Polink-2 Plus AP Broad Bulk kit	D68-110	110ml	(No cross react to RAT)	D65-110	110ml
			Polink-2 Plus AP Mouse-NR		
Polink-2 Plus AP Broad 18ml kit / 6ml kit	D68-18 / D68-6	18ml / 6ml	18ml/6ml kit (No cross react to RAT)	D65-18 / D65-6	18ml / 6ml
Polink-2 Plus AP Mouse bulk kit	D69-110	110ml	Fast Red Kit	C03-60	12 Tab + 60ml
Polink-2 Plus AP Mouse 18ml / 6ml kit	D69-18 / D70-6	18ml / 6ml	AP-Red+ Kit (40x concentrate)	C04-8	8ml
Polink-2 Plus AP Goat bulk	D66-110	110ml	BCIP/NBT Kit	C05-100/C05-18	100ml / 18ml
Polink-2 Plus AP Goat 18ml / 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 / 6ml kit	D67-18 / D67-6	18ml / 6ml			

Precautious: Please wear gloves and take other necessary precautions.

**Remarks:** For research use only.

22310 20th Ave Se #100, Bothell, WA 98021, USA Tel: 425.398.1500 Fax: 425.398.1519 http://www.gbi-inc.com