

Polink TS-MMR-Hu A Kit

(Polymer-HRP & AP triple staining kit)

(Detects two mouse and one rabbit primary antibodies on human tissue with DAB(Brown), GBI-Permanent Red (Red), and Emerald (Green))

Storage: 2-8°C

Catalog No.: TS301A-6 6mL* 60 slides**
 TS301A-18 18mL* 180 slides**
 TS301A-60 60mL* 600 slides**
**Volume of polymer conjugates*
*** If using 100µL per slide*

Intended Use:

The **Polink TS-MMR-Hu A Kit** is designed to use with user supplied two mouse primary antibodies and one rabbit primary antibody to detect three distinct antigens on a single human tissue or cell samples. TS-MMR-Hu has been tested on paraffin embedded tissue only; however, it may be used on frozen or freshly prepared monolayer cell smears. Please read through entire protocol as this protocol requires many steps to be done in their defined order. Choose from protocol TS301A protocol-1, TS301A protocol-2 and TS301A protocol-3 to accommodate your primary antibodies sensitivity to pre-treatment.

Triple staining uses traditional and non-traditional methods in immunohistostaining to reveal three distinct antigens and their co-expression on a single tissue^{1, 2}. **Polink TS-MMR-Hu A Kit** from GBI Labs (Golden Bridge International) supplies polymer enzyme conjugates: Polymer-HRP anti-rabbit, Polymer-AP anti-mouse and Polymer-HRP anti-mouse with three chromogens, DAB (brown); GBI-Permanent Red (red); and Emerald (green). **Polink TS-MMR-Hu A Kit** is a non-biotin system, avoiding non-specific binding caused by endogenous biotin. This kit has been optimized to have no cross detection when detecting two primary antibodies from the same host species using our unique blocking system. Simplified steps allow users to complete triple staining within 5 hours (without antigen retrieval) or 6 hours (with antigen retrieval). The well tested protocol provides user to permanently mount slides with coverslip.

Kit Components:

Component No.	Content	TS301A-6	TS301A-18	TS301A-60
Reagent 1	Mouse AP Polymer (RTU)	6mL	18mL	60mL
Reagent 2	Rabbit HRP Polymer (RTU)	6mL	18mL	60mL
Reagent 3A	DAB Substrate (RTU)	12mL	36mL	120mL
Reagent 3B	DAB Chromogen (20x)	1.5mL	2mL	6mL
Reagent 4A	GBI-Permanent Red Substrate (RTU)	15mL	36mL	120mL
Reagent 4B	GBI-Permanent Red Activator (5x)	3mL	7.2mL	24mL
Reagent 4C	GBI-Permanent Red Chromogen (100x)	150µL	360µL	1.2mL
Reagent 5	Antibody Blocker (40x)	30mL	100mL	100mLx3
Reagent 6A	TS-MMR Blocker A (RTU)	6mL	18mL	60mL
Reagent 6B	TS-MMR Blocker B (RTU)	6mL	18mL	60mL
Reagent 7	Mouse HRP Polymer (RTU)	6mL	18mL	60mL
Reagent 8	Emerald Chromogen (RTU)	6mL	18mL	60mL
Reagent 9	U-Mount (RTU)	6mL	18mL	NA

Protocol Notes:

- Proper Fixation: To ensure the quality of the staining and obtain reproducible performance, user needs to supply appropriately fixed tissue and well-prepared slides.
- Tissue needs to be adhered to the slide tightly to avoid falling off.
- Paraffin embedded sections must be deparaffinize with xylene and rehydrated with a graded series of alcohols before staining.
- Cell smear samples should be prepared as close to a monolayer as possible to obtain satisfactory results.
- Control slides are recommended for interpretation of results: positive, reagent (slides treated with Isotype control reagent), and negative control.
- DO NOT** let specimen or tissue dry during protocol. This will generate false positive and/or false negative signal.
- Important:** Never combine two antibodies from the same host species in one incubation step. Incubate 1st primary mouse antibody with rabbit antibody.
- 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.

Staining protocol selection and limitation of the kit:

- Most antigens will not be destroyed by heat. However, users need to check if there are proteins on the tissue that are heat sensitive before proceeding with the staining.
- TS301A Protocol-2 worksheet is suitable for one Mouse & one Rabbit primary Abs need pre-treatment, the other Mouse primary Ab is sensitive to pre-treatment.
- TS301A Protocol-3 worksheet is suitable when one Mouse & one primary antibody is sensitive to pre-treatment, but the second Mouse primary antibody needs pre-treatment.
- Please read the following table carefully before you start the experiment to ensure the result.
- This kit is not suitable for the following condition: 2 proteins are heat sensitive and detected by 2 mouse antibodies and one rabbit antibody requires HIER.

TS301A Staining Protocol 1:

Steps / Reagent	Staining Protocol	Incubation Time
1. Peroxidase and phosphatase Blocking Reagent: Supplied by user	a. Incubate slides in peroxidase and alkaline phosphatase blocking reagent (Klear Dual Enzyme Block E36 is Recommended) for 10 minutes. b. Rinse the slide using distilled water at least twice.	10 min
2. Antigen retrieval (optional): Refer to primary antibody data sheet	Note: Investigator needs to do antigen retrieval only one time during protocol see staining protocol a. Refer to primary antibody data sheet for antigen retrieval methods b. Wash with PBS/ 0.05% Tween20 for 2 minutes, 3 times.	
3. Primary Antibody Mix: Mix one Mouse and one Rabbit primary antibody: Supplied by user	Note: Investigator needs to optimize dilution prior to triple staining. DO NOT combine the same host species primary antibodies together at this step. a. Apply 2 drops or enough volume of mouse and rabbit primary antibody mixture to cover the tissue completely. Incubate in moist chamber for 30-60min. Recommend 30min to shorten total protocol time. b. Wash slides with PBS/ 0.05% Tween20 for 2 minutes, 3 times.	30 min
4. Reagent 1: Mouse AP Polymer (RTU)	a. Apply 1 to 2 drops (50-100µL) of Reagent 1 to cover the tissue completely. b. Incubate in moist chamber for 15-30 min. c. Wash slides with PBS/ 0.05% Tween 20 for 2 minutes, 3 times.	15-30 min
5. Reagent 2: Rabbit HRP Polymer (RTU)	a. Apply 1 to 2 drops of Reagent 2 to cover the tissue completely. b. Incubate in moist chamber for 15-30 min. c. Wash slides with PBS/ 0.05% Tween 20 for 2 minutes, 3 times.	15-30 min
6. Reagents 3A, 3B: Reagent 3A: DAB Substrate (RTU) Reagent 3B: DAB Chromogen (20x)	Note: Make enough DAB mix by adding 1 drop of Reagent 3B (DAB Chromogen) in 1mL of Reagent 3A (DAB Substrate). Mix well. Use within 7 hours. a. Apply 1 to 2 drops (50-100µL) of your DAB mixture to cover the tissue completely. b. Incubate for 5min. c. Rinse slides in multiple changes of distilled water 3 times, 2 min each time or under running tap water for 2minute.	5 min
7. Reagents 4A, 4B, 4C: Reagent 4A: GBI-Permanent Red Substrate (RTU) Reagent 4B: GBI-Permanent Red Activator (5x) Reagent 4C: GBI-Permanent Red Chromogen (100x)	Note: Shake GBI-Permanent Red Activator before adding into GBI-Permanent Red Substrate . a. Add 200µL of Reagent 4B (Activator) into 1mL of Reagent 4A (Substrate) and mix until clear. Add 12µL of Reagent 4C (Chromogen) into the mixture and mix well. [Note: For fewer slides, add 100µL of Reagent 4B (Activator) into 500µL of Reagent 4A (Substrate buffer) and mix until clear. Add 6µL of Reagent 4C (Chromogen) into the mixture and mix well]. 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.	10 min
8. Reagent 5: Antibody Blocker (40x)	Note: This step will block antibodies of previous step so no cross reaction will occur in this protocol. HIER can be done immediately after Antibody Blocker step if the primary antibodies require antigen retrieval. For frozen tissues, a lower temperature of 65°C must be used during the Antibody Blocker step to prevent dissociation of the tissue from the slide. a. Use hot plate or water bath to heat diluted Reagent 5 (Antibody Blocker) to 1x solution (1 part of Antibody Blocker in 39 parts of distilled water) to 80°C. Make enough volume to cover the tissue in beaker. b. Put slides in heated Antibody Blocker for 10 minutes at 80°C. c. Remove slides from the Antibody blocker; cool slides 5 seconds. d. Rinse slides in multiple changes of distilled water. If antigen retrieval step is required, go directly to step 8 if not complete step 7e and move on to step 9 . e. Wash with PBS/ 0.05% Tween20 for 2 minutes, 3 times.	10 min

9. Antigen retrieval: Refer to primary antibody data sheet	a. Refer to primary antibody data sheet for antigen retrieval methods. b. Wash slides with PBS/ 0.05% Tween20 for 2 minutes, 3 times.	Up to 1 hour
10. Reagent 6A: TS-MMR Blocker A (RTU)	a. Apply 2 drops or enough volume of Reagent 6A (DS-MMR Blocker A) to cover the tissue completely. Mix well on the slide and incubate in moist chamber for 30 min. b. Rinse with PBS containing 0.05% Tween-20 for 2 min., 3 times.	30 min
11. Reagent 6B: TS-MMR Blocker B (RTU)	a. Apply 2 drops or enough volume of Reagent 6B (DS-MMR Blocker B) to cover the tissue completely. Mix well on the slide and incubate in moist chamber for 5 min. b. Rinse with PBS containing 0.05% Tween-20 for 2 min., 3 times.	5 min
12. 2 nd Mouse primary antibody: Supplied by user	Note: Investigator needs to optimize dilution prior to triple staining. a. Apply 2 drops or enough volume of the 2 nd mouse primary antibody to cover the tissue completely. Incubate in moist chamber for 30-60 min. Recommend 30 minutes to shorten total protocol time. b. Wash slides with PBS/ 0.05% Tween-20 for 2 minutes, 3 times.	30 min
13. Reagent 7: Mouse HRP Polymer (RTU)	a. Apply 1 to 2 drops (50-100µL) of Reagent 7 (Mouse HRP Polymer) to cover the tissue completely. Incubate slides in moist chamber for 15-30 min. b. Wash slides with PBS/ 0.05% Tween20 for 2 minutes, 3 times.	15-30 min
14. Counterstain: (Optional but must be done before Emerald Chromogen step): Not provided	Note: If two antigens are co-localized in the nucleus, you want less counter stain to optimize the visualization in the nucleus; however, you can counter stain using normal protocol time if antigens are co-localized in cytoplasm or membrane or the three antigens are localized in different cells. a. Counterstain dip in diluted hematoxylin for 5 seconds for nuclear co- localization or 30 seconds for cytoplasmic or membrane co- localization. DO NOT over stain with hematoxylin. b. Rinse thoroughly with tap water for 1min. c. Put slides in PBS for 5-10 seconds to blue, DO NOT over blue. d. Rinse well in distilled or tap water for 1min. e. Wash slides with PBS/ 0.05% Tween20 for 2 minutes, 3 times.	5 seconds
15. Reagent 8: Emerald Chromogen (RTU) Do hematoxylin first	a. Apply 1 to 2 drops (50-100µL) of Reagent 8 (Emerald Chromogen) to cover the tissue completely. b. Incubate slides in humid chamber for 5 minutes. c. Wash slides in tap water for 3 times for 30 seconds! Important: Emerald Chromogen is water soluble, do counter stain first. Do not leave slides sitting in water. Always stain Emerald chromogen AFTER GBI-Permanent Red stain because GBI-Permanent Red removes the Emerald and after hematoxylin.	5 minutes
16. Dehydrate section: It is important to follow the protocol	Note: Please wipe off extra water and air-dry slides before dehydration and clear. a. Dehydrate with 85% ethanol 20 sec b. Dehydrate with 95% ethanol 20 sec c. Dehydrate with 100% ethanol 20 sec d. Dehydrate with 100% ethanol 20 sec e. Dehydrate with 100% ethanol 20 sec f. Dehydrate with xylene 20 sec CAUTION: DO NOT dehydrate in xylene longer than 20 seconds! It will erase GBI-Permanent Red stain!	2 min
17. Reagent 9 U-Mount (RTU)	a. Apply 1 drop (50µL) of Reagent 9 (U-Mount) to cover the tissue section and apply glass coverslip. b. Apply force to coverslip to squeeze out any extra mountant and bubbles for optimal clarity. Removing excess also to prevent leaching of GBI-Permanent Red stain.	

Troubleshooting

Problem	Tips
Uneven stain on 3 primary antibodies	1. Need to adjust the titer of each antibody. 2. The amount of each protein expressed on tissue may be different. 3. Set slides in water too long so that Emerald is washed away. 4. Set slides in Xylene too long so that GBI-Permanent Red is washed away.
Emerald Chromogen is blue not green when non-co-localized with GBI Permanent Red.	1. Emerald should be green when non colocalized with GBI-Permanent Red. If Emerald chromogen is blue the titer on the primary antibody is not dilute enough for the protocol. Re-titer primary antibodies individually first.
No stain on 1 or 2 antibodies	1. Missing steps or steps reversed.
Green Background on the slide	1. Titer primary antibody.
GBI-Permanent Red is leaching	1. Use fresh 100% ethanol and xylene. 2. Slide sat too long in xylene. Do not go over 20seconds!
Artifacts on slides	1. Slides not completely dried before mount. Use fresh 100% Ethanol and xylene. 2. GBI-Permanent Red Activator was not properly mixed into the Substrate. Mix until clear.

Precautions:

Please wear gloves, eye protection, and take other necessary precautions. If any of the reagents come into contact with skin, wash area completely with plenty of water and soap. If irritation develops seek medical attention.

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 TS301A 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 ensure 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 “√” each step during the experiment
- Steps follow de-paraffinization
- Refer to insert for details of each step

TS301A Protocol-1 is suitable when all primary antibodies need pre-treatment, or all primary antibodies do not need pre- treatment.

	Main Protocol Step	TS301A Protocol-1	Experiment 1 Date:	Experiment 2 Date:	Experiment 3 Date:	Experiment 4 Date:
1	Step 1	Peroxidase or Alkaline Phosphatase Block E36 is recommended. User supplied User supplied				
2	Step 2	HIER(Optional)				
3	Step 3	Mouse 1°Ab & Rabbit 1°Ab mix User supplied (30-60 min.)				
4	Step 4	Reagent 1 Mouse AP Polymer (15-30 min.)				
	Step 5	Reagent 2 Rabbit HRP Polymer (15-30 min.)				
5	Step 6	Reagent 3A & Reagent 3B DAB requires mixing (5 min.)				
6	Step 7	Reagent 4A & Reagent 4B GBI-Permanent Red requires mixing (10 min)				
7	Step 8	Reagent 5 Antibody Blocker (10min)				
8	Step 10	Reagent 6A DS-MMR Blocker A RTU (30min)				
9	Step 11	Reagent 6B DS-MMR Blocker B RTU (5min)				
10	Step 12	Mouse 1°Ab User supplied (30-60 min)				
11	Step 13	Reagent 7 Mouse HRP Polymer RTU (15 min)				
12	Step 14	Counter stain (Note 2) User supplied (5-10 sec)				
13	Step 15	Reagent 8 Emerald Chromogen RTU (5min)				
14	Step 16	It is important to follow the protocol. To maintain stain! Dehydrate section 20seconds for each step				
15	Step 17	Reagent 9 U-Mount RTU Mount & coverslip				
	Result	Stain pattern on controls is correct: Fill in Yes or NO				

Note1: Normal wash steps = Wash with PBS containing 0.05% Tween-20 for 3 times for 2 min each.

Note2: *Using as a co-localization staining kit,

If antigens are co-localized in nucleus counter stain and blue should be for 5 seconds to blue.

If antigens are co-localized in cytoplasm and membrane or in different cells counter stain using normal protocol time.

TS301A Protocol-2 is suitable when one Mouse & one Rabbit primary antibodies need pre-treatment, but the second Mouse primary antibodies is sensitive to pre-treatment.

	Main Protocol Step	TS301A Protocol-2	Experiment 1 Date:	Experiment 2 Date:	Experiment 3 Date:	Experiment 4 Date:
1	Step 1	Peroxidase or Alkaline Phosphatase Block E36 is recommended. User supplied User supplied				
2	Step 12	Mouse 1°Ab (sensitive to HIER) User supplied (30-60min)				
3	Step 13	Reagent 7 (RTU) Mouse HRP Polymer RTU (15min)				
4	Step 6	Reagent 3A&3B DAB requires mixing (5 min)				
5	Step 8	Reagent 5 Antibody Blocker require mixing (10min)				
6	Step 9	HIER (DAB will not be removed)				
7	Step 10	Reagent 6A (RTU) DS-MMR Blocker A RTU (30min)				
8	Step 11	Reagent 6B (RTU) DS-MMR Blocker B RTU (5min)				
9	Step 3	Mouse 1°Ab & Rabbit 1°Ab mix (Abs requires HIER) User supplied (30-60 min)				
10	Step 4 & 5	Reagent 1 & Reagent 2 Mouse AP Polymer & Rabbit HRP Polymer (15-30 min each)				
11	Step 7	Reagent 4A & Reagent 4B GBI-Permanent Red requires mixing (10 min)				
12	Step 14	Counter stain (Note 2) User supplied (5-10 sec.)				
13	Step 15	Reagent 8 Emerald Chromogen RTU (5min.)				
14	Step 16	It is important to follow the protocol. To maintain stain! Dehydrate section 20seconds for each step				
15	Step 17	Reagent 9 U-Mount RTU Mount & coverslip				
	Result	Stain pattern on controls is correct: Fill in Yes or NO				

Note1: Normal wash steps = Wash with PBS containing 0.05% Tween-20 for 3 times for 2 min each.

Note2: *Using as a co-localization staining kit,

If antigens are co-localized in nucleus counter stain and blue should be for 5 seconds to blue.

If antigens are co-localized in cytoplasm and membrane or in different cells counter stain using normal protocol time.

TS301A Protocol-3 is suitable when one Mouse & one Rabbit primary antibodies are sensitive to pre-treatment, but the second Mouse primary antibody needs pre-treatment.

	Main Protocol Step	TS301A Protocol-3	Experiment 1 Date:	Experiment 2 Date:	Experiment 3 Date:	Experiment 4 Date:
1	Step 1	Peroxidase or Alkaline Phosphatase Block E36 is recommended. User supplied User supplied				
2	Step 3	Mouse 1°Ab & Rabbit 1°Ab mix User supplied (30-60 min.)				
3	Step 4 & 5	Reagent 1 & Reagent 2 Mouse AP Polymer & Rabbit HRP Polymer (15-30 min each)				
4	Step 6	Reagent 3A&Reagent 3B DAB require mixing (5 min)				
5	Step 7	Reagent 4A&Reagent 4B GBI-Permanent Red requires mixing (10min)				
6	Step 8	Reagent 5 Antibody Blocker (10min)				
7	Step 2	HIER				
8	Step 10	Reagent 6A DS-MMR Blocker A RTU (30min)				
9	Step 11	Reagent 6B DS-MMR Blocker B RTU (5min)				
10	Step 12	Mouse 1°Ab (sensitive to HIER) User supplied (30-60 min.)				
11	Step 11	Reagent 7 (RTU) Mouse HRP Polymer (15 min.)				
12	Step 14	Counter stain (Note2) User supplied				
13	Step 15	Reagent 8 (RTU) Emerald Chromogen (5min)				
14	Step 16	It is important to follow the protocol to maintain stain! Dehydrate section 20seconds for each step				
15	Step 17	Reagent 9 U-Mount RTU Mount & coverslip				
	Result	Stain pattern on controls is correct: Fill in Yes or NO				

Note1: Normal wash steps = Wash with PBS containing 0.05% Tween-20 for 3 times for 2 min each.

Note2: *Using as a co-localization staining kit,

If antigens are co-localized in nuclear counter stain and blue should be for 5 seconds to blue.

If antigens are co-localized in cytoplasm and membrane or in different cells counter stain using normal protocol time.