

7-Color Manual IHC Kit (pH9.0)

[Product No.]
OM642683

[Principle of action] The 7 color IHC detection kit, get rid of the traditional immunofluorescence experimental conditions of antibody species sources of restrictions and constraints, without ventilation kitchen and traditional dewaxing repair tank, the traditional three times xylene dewaxing, three to five times gradient ethanol hydration and antigen repair integration into a solution, DAIP and sealing integration into a solution, effectively shorten the Three fluorescent dyes immunohistochemical experiments used in the reagent one-stop ready, with high sensitivity of HRP polymer secondary antibody (Anti-Rabbit/Mouse) and fluorescence color solution supporting the use, can be a color seven antibodies (without considering the source of antibody species), making the kit has the characteristics of convenient operation, rapid, high sensitivity.

[Packaging Specifications] 1mL 2mL 5mL

[Storage conditions and validity]

- 1, storage requirements: see the table.
- 2, validity: one year

[Self-provided materials]

- 1, suitable heating device, dye vat, pipette, cover glass and other commonly used consumables
- 2, anhydrous alcohol
- 3, neutral gum
- 4, PBST, pure water and other commonly used reagents

[Components in kit]

Reagent	Name	Size			Storage
		1mL	2mL	5mL	
Reagent 9A	Dewaxing and Repair 20X	60 mL	60 mL	200mL	2~8°C
Reagent B	Blocking solution	1mL	2mL	5mL	2~8°C
Reagent C	Polymer HRP Mouose/Rabbit	6mL	6mL*2	30mL	2~8°C
Reagent D-425	425 labeled Tyramide	1mL	2mL	5mL	2~8°C
Reagent D-488	488 labeled Tyramide	1mL	2mL	5mL	2~8°C
Reagent D-525	525 labeled Tyramide	1mL	2mL	5mL	2~8°C
Reagent D-594	594 labeled Tyramide	1mL	2mL	5mL	2~8°C
Reagent D-680	680 labeled Tyramide	1mL	2mL	5mL	2~8°C
Reagent	750 labeled Tyramide	1mL	2mL	5mL	2~8°C

D-750					
Reagent G	DAPI/mounting solution	1mL	2mL	5mL	-20°C
Reagent F	Stripping solution 20X	60mL	60mL	200mL	2~8°C

[Intended Use]

Immunohistochemical fluorescence staining, with Immunoway conventional rabbit/mouse antibodies or ready-to-use antibody.

[Sample requirements]

Paraffin or frozen sections, cell slides. Recommended with Immunoway conventional concentrated or ready-to-use primary antibody.

Paraffin sections: primary antibody application indicated IHC-p

Frozen sections: primary antibody application indicated IHC-f


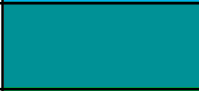



Cell slides: primary antibody application indicated IF or ICC.






Some frozen or tissue sections are easy to fall off after repeated heating, the tissue can be put into the 24/12/6 well plate.

[Read before use]

- 1, Use deionized water to dilute reagent 9A, reagent F, volume ratio 1:19.
- 2, If you need to send or vertical preservation slices, please drop nail polish or neutral gum around the cover glass to dry.
- 3, Paraffin slices need to be put into the oven at 65 °C for one hour before use. Freezing slices, cell climbing slices have no this step.
- 4, TSA staining is more sensitive than conventional fluorescence staining and TMB staining. If the fluorescence intensity is too high, you can try to double dilute of primary antibody, or double dilute of Tyramide reagent.
- 5, The expression abundance of indicators can be determined by conventional staining or pre-experiment. Usually, the highest expression index is stained first, and the lowest expression index is stained last.
- 6, 525, 555, 594 markers are different degrees red under the microscope. Please use special imaging instruments and analytical software to distinguish the spectra.
- 7, The dewaxing antigen repair solution in this kit is suitable for most primary antibodies. If the staining effect is not good, you can use the general public formula to dewax the slices and repair antigen.
- 8, Because of the small adhesion ability of frozen slices and cell climbing slices, the pre-experiment of **step 9** can be carried out separately before use this kit to ensure that the phenomenon of dropping slices does not occur.

[Recommended fluorescence channel for imaging]

Reagent	Nmae	Exi	Emi	Filter	Color
Reagent D-350	350 labeled Tyramide	350	450	DAPI	
Reagent D-435	435 labeled Tyramide	435	470	Aqua,Opal470	
Reagent D-488	488 labeled Tyramide	496	519	FITC, Opal520	
Reagent D-525	525 labeled Tyramide	530	560	Cy3,SpGold,Opal540	
Reagent D-555	555 labeled Tyramide	555	567	TRITC, Cy3,SpOr,Opal570	

Reagent D-594	594 labeled Tyramide	590	617	SpRed,TxRed,Opal620	
Reagent D-647	647 labeled Tyramide	650	665	Cy5,Opal650	
Reagent D-680	680 labeled Tyramide	680	700	Cy5.5,Opal690	
Reagent D-750	750 labeled Tyramide	750	796	Cy7,Opal780	
Reagent G	DAPI/mounting solution	345	455	DAPI	

* Due to the overlap of excitation and emission spectra, Opal540/570 can be observed mixed under fluorescence microscope objective. Cy5/Cy5.5 can be observed mixed

** Using the full-spectrum imaging system can split separate spectra to achieve more than 6 colors imaging

[Procedure for paraffin sections]

1. Put reagent 9A (dewaxing repair 2-in-1) working solution into the repair box and heat it to boiling.
2. Put the section into the boiling reagent 9A (dewaxing repair 2-in-1) working solution (to ensure the pH value of the repair solution, metal section holder cannot be used), and the liquid completely immerses the tissue on the section. Heat it continuously for 30 min.

Note: Antigen repair is depended on a variety of factors such as the primary antibody supplier, tissue type, etc., and the commonly used antigen repair reagents or other validated repair conditions in the laboratory can be used. This step can be omitted for tissues with frozen slides or cell-crawling slides which are easy to fall off.

3. Remove the repair box from the heating source and cool it to room temperature.
4. Take out the section into a beaker filled with distilled water, and then soak it with distilled water 5-6 times.
5. Drain the slide for a few seconds, wipe the water around the tissue with filter paper, drop reagent B (peroxidase blocking buffer) 50-100ul (it can cover the tissue), incubate at room temperature for 15 min, and wash it with PBST for 2 min × 3 times.
6. Drain the slide for a few seconds, wipe off the moisture around the tissue with a filter paper, circle the tissue with an immunohistochemical pen (end to end should be closed, and cannot be drawn on the tissue), dilute the primary antibody, and drop the corresponding primary antibody onto the tissue until completely covered. Allow to incubate at room temperature or 37 °C for 1 to 2 hours, or incubate at 37 °C for 30 minutes after overnight at 4 °C in a wet box, and rinse with PBST for 2 minutes × 3 times.
7. Drain the slide for a few seconds, wipe off the moisture around the tissue with a filter paper, drop reagent C (HRP polyclonal anti-rabbit/mouse secondary antibody) working solution 50-100ul (to cover the tissue), incubate at room temperature for 30 minutes, and rinse with PBST for 2 minutes × 3 times.
8. Add fluorescent reagent D-594 working solution 50-100ul (to cover the tissue), after 10 minutes, rinse with PBST for 2 minutes × 3 times.
9. Place the slices in the repair box, add reagent F, and microwave for 3 minutes on high power and 15 minutes on low power. After cooling naturally, rinse with PBST for 2 minutes × 3 times. (Because of different microwave powers, the time may need to be adjusted, and maintain the liquid above 95 °C for 15 minutes)
10. Repeat steps 6-9, and change the fluorescent dye to reagent D-488, and incubate the second primary antibody.
11. Repeat steps 6-9, and change the fluorescent dye to reagent D-425, and incubate the 3th primary antibody.
12. Repeat steps 6-9, and change the fluorescent dye to reagent D-680, and incubate the 4th primary antibody.
13. Repeat steps 6-9, and change the fluorescent dye to reagent D-525, and incubate the 5th primary antibody.
14. Repeat steps 6-8, and change the fluorescent dye to reagent D-750, and incubate the 6th primary antibody.

15. Drop a drop of reagent G (about 30-50ul) on the tissue slice, cover the cover glass, let the slice contact the sealing solution, try to avoid bubbles, scan or take pictures

Note: The above fluorescence sequence is applicable to situations where the expression levels of various indicators are not significantly different. The fluorescence dye sequence can be adjusted based on the expression of corresponding indicators and the results of preliminary experiments. For example, high-expression indicators can be labeled preferentially, followed by low-expression indicators. Some tissues may be affected by multiple rounds of heating, which may affect the binding of primary antibodies to proteins. It is recommended to adjust the staining sequence through multiple rounds of heating with a single color.

[Procedure for cell slices/frozen sections]

1. **Optional step:** fixation: after restoring the section to room temperature, use 4% paraformaldehyde (not provided in the kit) for 10-15min, wash with 1xPBS three times, 5 minutes each time. This step is usually added in the cell crawling slices. The frozen slices are not fixed before OCT embedding. It is recommended to add this step.

2. **Optional step:** antigen repair: reagent 9A (dewaxing repair two-in-one) working solution is put into the repair box, heated to boiling. The slides were placed in boiling reagent 9A (dewaxing repair 2-in-1) working solution (to ensure the pH value of the repair solution, metal slide holders were not used), and the liquid completely immersed the tissue on the slides. The repair box was removed from the heating source and cooled to room temperature.

Note: antigen repair is depended on a variety of factors such as the primary antibody supplier, tissue type, etc., and the commonly used antigen repair reagents or other validated repair conditions in the laboratory can be used. This step can be omitted for tissues with frozen slides that are easy to fall off, or for cell-crawling slides or water bath repair (see Step 9).

3. **Optional steps:** permeate with 0.5% Triton X-100 (not provided in the kit) at room temperature for 20 min (this step is omitted for antigens expressed on the cell membrane).

4. Wash with 1xPBS three times, 2 minutes each time.

5. Drain the slide for a few seconds, wipe off the moisture around the tissue with filter paper, drop reagent B (peroxidase blocking buffer) 50-100ul (enough to cover the tissue), incubate at room temperature for 15 min, and wash with PBST for 2 min × 3 times.

6. Drain the slide for a few seconds, wipe off the moisture around the tissue with filter paper, circle the tissue with an immunohistochemical pen (the ends should be closed, and cannot be drawn onto the tissue), dilute the primary antibody, and drop the corresponding primary antibody onto the tissue until completely covered. After incubation at room temperature or 37 °C for 1 to 2 hours, or after incubation at 4°C overnight and rewarming at 37 °C for 30 minutes, PBST washed 2 min × 3 times.

7. Drain the slide for several seconds, wipe off the moisture around the tissue with filter paper, drop reagent C (HRP polyclonal anti-rabbit/mouse secondary antibody) working solution 50-100ul (can cover the tissue), incubate at room temperature for 30 minutes, PBST washed 2 min × 3 times.

8. Add fluorescent reagent D-594 working solution 50-100ul (can cover the tissue), 10 minutes later, PBST washed 2 min × 3 times.

9. Place the slices in the repair box and add reagent F. Place the repair box in a water bath containing boiling distilled water and heat for 20 minutes.

10. Repeat steps 6-9, and change the fluorescent dye to reagent D-488, and incubate the second primary antibody.

11. Repeat steps 6-9, and change the fluorescent dye to reagent D-425, and incubate the 3th primary antibody.

12. Repeat steps 6-9, and change the fluorescent dye to reagent D-680, and incubate the 4th primary antibody.

13. Repeat steps 6-9, and change the fluorescent dye to reagent D-525, and incubate the 5th primary antibody.

14. Repeat steps 6-8, and change the fluorescent dye to reagent D-750, and incubate the 6th primary antibody.

15. Drop a drop of reagent G (about 30-50ul) on the tissue slice, cover the cover glass, let the slice contact the sealing solution, try to avoid bubbles, scan or take pictures

Note: The above fluorescence sequence is applicable to situations where the expression levels of various indicators are

not significantly different. The fluorescence dye sequence can be adjusted based on the expression of corresponding indicators and the results of preliminary experiments. For example, high-expression indicators can be labeled preferentially, followed by low-expression indicators. Some tissues may be affected by multiple rounds of heating, which may affect the binding of primary antibodies to proteins. It is recommended to adjust the staining sequence through multiple rounds of heating with a single color.

[Product Performance Indicators]

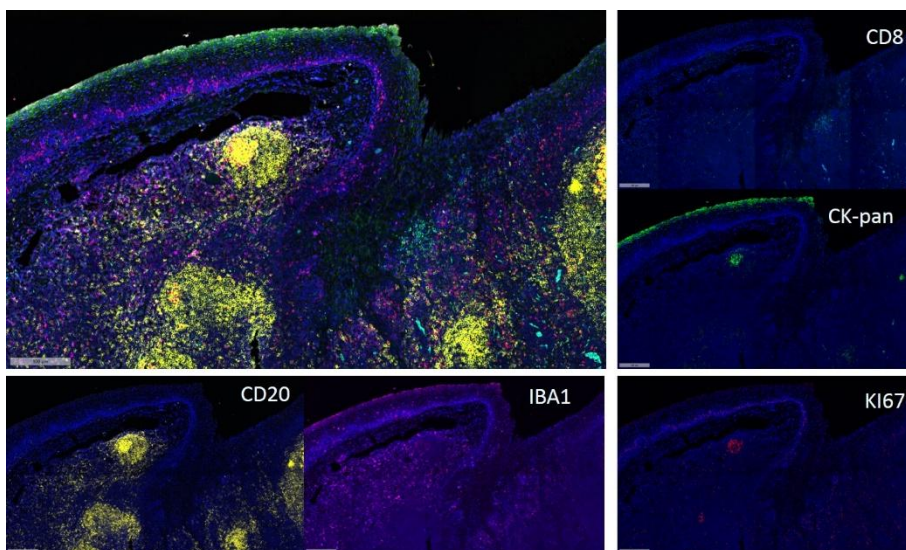
1. Compliance: The positive control result is positive, and the location of positive staining should be accurate, without background staining; the blank control and negative control staining results are negative.
2. Intra-batch repeatability: There is no significant difference in the intensity and location of staining in tissue sections from the same tissue source.
3. Inter-batch repeatability: There is no significant difference in the intensity and localization of staining between different batches of reagents on tissue sections of the same tissue origin.

[Precautions]

1. This product is only for scientific research and is not for other purposes.
2. Requires professional use.
3. Apply appropriate protective measures to avoid contact between reagents and the skin and eyes.
4. Waste liquid treatment: Conduct harmless treatment and comply with relevant environmental protection requirements.
5. The operation process needs to keep the wet tissue of the slide, such as dry pieces, will lead to non-specific staining results.
6. In the process of continuous heating repair, using small fire to maintain boiling, do not heat the dewaxing repair reagent splash out of the beaker.

[Frequently Asked Questions]

- 1, deep staining: a high concentration of anti-high, too long time.
- 2, light or no staining: a low concentration of anti-low, too short time.
- 3, no specific staining: slice dewaxing is not complete. Can be appropriately prolonged baking time.



Multiplex Fluorescence immunohistochemical analysis of Human tonsil tissue (formalin-fixed paraffin-embedded section). CK-pan mouse mAb(YM6815 Immunoway) green, Ki-67 rabbit mAb(YM7002 Immunoway) red, Iba 1 mouse

mAb(YM4765 Immunoway) purple, CD8 a mouse mAb(YM4815 Immunoway) cyan, CD20 mouse mAb(YM4814 Immunoway) yellow, The section was incubated in 5 rounds of staining; sequentially for Anti-antibodies; each using a separate fluorescent tyramide signal amplification system. EDTA based antigen retrieval (Immunoway YS0004, pH 9.0, 20 minutes) was used in between rounds of tyramide signal amplification to remove the antibody from the previous round, to avoid any cross-reactivity. DAPI (dark blue) was used as a nuclear counter stain. Microscopy and pseudocoloring of individual dyes was performed using a Slideviewer Imaging System (Excilone).

[Related product]

Catalog	Product Name	Size
OM642680	4-Color Manual IHC Kit	1mL 2mL 5mL
OM642679	3-Color Manual IHC Kit	1mL 2mL 5mL
OM642681	5-Color Manual IHC Kit	1mL 2mL 5mL
OM642682	6-Color Manual IHC Kit	1mL 2mL 5mL
OM642683	7-Color Manual IHC Kit	1mL 2mL 5mL

[Individual components]

Catalog	Reagent	Size
OM642680	Polymer HRP Goat Anti Mouse/Rabbit IgG(H+L)	3mL 10mL 100mL
OM642679	350 labeled Tyramide	2mL 5mL 10mL
OM642681	435 labeled Tyramide	2mL 5mL 10mL
OM642682	488 labeled Tyramide	2mL 5mL 10mL
OM642683	525 labeled Tyramide	2mL 5mL 10mL
OM642680	555 labeled Tyramide	2mL 5mL 10mL
OM642679	594 labeled Tyramide	2mL 5mL 10mL
OM642681	647 labeled Tyramide	2mL 5mL 10mL
OM642682	680 labeled Tyramide	2mL 5mL 10mL
OM642683	750 labeled Tyramide	2mL 5mL 10mL
OM642680	Antigen Repair 20X pH9.0	250mL
OM642679	Antigen Repair 20X pH6.0	250mL
OM642681	DAPI/mounting solution	5mL
OM642682	Stripping solution 20X	200mL