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RNAscope for FFPE Mouse Tissue

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We use this protocol and it's working

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Abstract

This protocol details the RNAscope for FFPE Mouse Tissue.



Materials

RNAscope Multiplex Fluorescent Reagent Kit v2 (Cat. #323100).

RNAscope® Multiplex Fluorescent Reagent Kit v2 Advanced Cell Diagnostics Catalog #323100

Day 1:

- Fresh Xylene
- Absolute alcohol
- Hydrogen peroxide & Protease plus from same kit
- Absorbent paper
- 10X Target retrieval
- C1, C2, C3 probes
- X RNAscope[™] 3-plex Negative Control Probe Advanced Cell Diagnostics Catalog #320871
- RNAscope™ 3-plex Positive Control Probe- Mm Advanced Cell Diagnostics Catalog #320881
- 1X wash buffer

Day 2:

- AMP1, AMP2, AMP3
- HRP-C# (need C1-C3 if developing all 3 channels)
- TSA buffer
- HRP-Blocker
- Opal 570 Reagent Pack Perkin Elmer Catalog #FP1488001KT
- 🔯 Opal 690 Reagent Pack **Perkin Elmer Catalog #**FP1497001KT
- TSA-DIG & Opal 780 (Cat. #FP1501001KT)
- 1X Antibody Diluent (Cat. #ARD1001EA)
- DAPI
- X TrueBlack® Plus Lipofuscin Autofluorescence Quencher, 40X in DMSO Biotium Catalog #23014
- ProLong™ Gold Antifade Mountant Invitrogen Thermo Fisher Catalog #P36930
- Cover Slip
- Tween 20
- PBS

Solutions

1X Wash Buffer

■ 40 mL 50X WashBuffer + 4 1960 mL RNase-free water



Target Retrieval

■ 🗸 25 mL 10X Target Retrieval + 🗸 225 mL RNase-free water

5x SSC

 \perp 50 mL 20X SSC + \perp 150 mL RNase-freewater

TSA-DIG: for 8 samples

 \perp 2 μ L TSA-DIG + \perp 1000 μ L TSA buffer

Opal 780 Dye: for 8 samples

 \perp 8 μ L opal 780 + \perp 1000 μ L 1X Antibody Diluent

1X TrueBlack Plus: for 8 samples

 Δ 25 μL 40X TrueBlack Plus + Δ 1000 μL 1X PBS \rightarrow **VORTEX**

probe channel	Channel 1 (C1)	Channel 2 (C2)	Channel 3 (C3)
channel sensitivity	highest	weakest	high
cell type analysis of target gene expression	gene of interest	cell type marker 1 (e.g. vGLUT1/2)	cell type marker 2 (e.g. GAD1/2)

Troubleshooting



DAY 1 - Bake/Adhere 1h Spray RNase away on bench top, slide holder, metal container. 2 Warm oven and slide holder up to \$\mathbb{\mod}\mathbb{\mod}\mathbb{\math}\m{\mathbb{\mathbb{\mathbb{\ 3 Label the lower part of the slide with a pencil. Can place slides on bench (since sprayed with RNase away). 4 Put slides into slide holder and place into oven \rightarrow bake for \bigcirc 01:00:00 \bigcirc 60 °C. 1h Meanwhile.. 4.1 Turn plate warmer on to \$\\\$\\\$ 60 \cdot \C \. 4.2 Make Target Retrieval solution. 4.3 Fill Xylene and Ethanol containers. Note After Baking: Possible stopping point: Store @ \$\ \mathbb{\mathbb{R}} \ \text{Room temperature} \, good for \(\sigma \) **(*)** 168:00:00 • 5 Set hyb oven temp to 40 °C. 5.1 Wet humidifying paper with nanopure water (does not need to be dripping). 5.2 Place paper on bottom of slide tray and put into the oven for 00:30:00 @ 30m \$ 40 °C ⋅

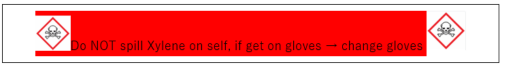
6 Place slides into tissue tek container and transport to fume hood.

DAY 1 - Deparaffinize

5m

7

5m



Xylene container 1 for 00:05:00.

- 7.1 Meanwhile.. Turn on vegetable steamer.
- 7.2 When transferring to container $2 \rightarrow$ shake off excess.
- 8 Xylene container 1 for 00:05:00 .

5m

- 8.1 When transferring to ethanol \rightarrow shake off excess.
- 9 Absolute ethanol container 1 for 00:02:00.

2m

- 9.1 When transferring to container $2 \rightarrow$ shake off excess.
- 10 Absolute ethanol container 2 for 00:02:00.

2m

11 Dry on slide warmer for ~ 👏 00:05:00 @ 🖁 60 °C until dry.

5m

11.1 **Meanwhile...** clean bench with RNase away.



DAY 1 - Hydrogen Peroxide

10m

- Place slides on bench top and cover tissue completely with Hydrogen Peroxide (~5-8 drops).
- Incubate for 00:10:00 @ Room temperature.



- Meanwhile..
- 1. Boil Target Retrieval solution and a container of Nano pure water in microwave until boiling.
- 13.1 *Microwave for (5) 00:01:00 at a time*



13.2 When reaches a boil \rightarrow place containers in warm steamer to ensure solution is at least $99 \, ^{\circ}\text{C}$.



- Remove solution using absorbent paper by tapping long side on paper.
 - Place in slide holder.
- 15 Wash slides in slide holder with \angle 200 mL of Nano pure water ~3-5 dunks.



Take slides out and repeat step **■** - **■** 1x

DAY 1 - Target Retrieval Step



- 17 Keep in water and transport to steamer.
- 17.1 Dunk a couple times and soak for 00:00:10 in nanopure water (steamer).

10s



- 18 Place in target retrieval solution in the steamer for 00:30:00 (for brain samples) or (for other tissue).
- 45m

19 Rinse slides in fresh \triangle 200 mL of Nano pure water \rightarrow 00:00:15.

15s

20 Transfer slides to the 2nd container of absolute ethanol in the fume hood \rightarrow **(:)** 00:03:00 .

3m

21 Dry slides in slide warmer - ~ (5) 00:05:00 @ # 60 °C.

5m

- Meanwhile...
- 21.1 Rinse slide holder in DI water in sink and let dry on paper towel.
- 21.2 Spray bench with RNase away.

DAY 1 - Barrier/Protease Plus



- 22 Put dry slides on bench and square off with hydrophobic pen.
- 22.1 Leave a little extra room at one side of square to allow space to aspirate later.
- 23 Apply ~5 drops of protease plus to tissue until sample is completely covered.
- 24 Incubate in oven for 00:30:00 @ \$ 40 °C .



- Meanwhile..
- 24.1 Warm probes @ **▮** 40 °C **for ~ ♦** 00:10:00 **.**



10m

- 24.2 Create probe solution.



Note

C1 probe is at $1X \rightarrow assign to low expresser gene$ C2 & C3 probes are at 50 X \rightarrow dilute w/ C1 probe if using C1 or with probe diluent to 1X

25 Wash with DI water in wash tray - 2X.

26 Aspirate with \triangle 200 μ L pipette tip in the fume hood.

26.1 Wipe bottom of wash tray with big kimwipe.

DAY 1 - Hybridize Probe



27 Add $\sim \Delta$ 120 µL of probe mix to respective samples \rightarrow cover sample completely.



28 Incubate for (5) 02:00:00 (a) \$\mathbb{8}\$ 40 °C .



29 Wash in 1X wash buffer for 2 minutes 2X.



Note

Store till day 2 in 5X SSC @ Broom temperature .

- Pour ethanol back into reagent bottle if not doing RNAscope within the next week so doesn't evaporate.
- 2m

29.1 Wash in 1X wash buffer for 600:02:00 (1/2).



29.2 Wash in 1X wash buffer for 600:02:00 (2/2).

2m





DAY 2

2m

- 30 Spray RNase away on bench top, slide holder, metal container.
- 31 Hydrate Paper and turn on oven to 4 40 °C.



Wash slides with 1X wash buffer for 2 minutes with slight agitation 2X.



Wash slides with 1X wash buffer for 00:02:00 with slight agitation (1/2).



32.2 Wash slides with 1X wash buffer for 00:02:00 with slight agitation (2/2).



- 33 Aspirate.
- 34 Hybridize AMP1.
- 34.1 4-5 drops covering the sample with AMP1.
- 34.2 Incubate for 00:30:00 @ \$ 40 °C .



34.3 Wash slides with 1X wash buffer for 2 minutes with slight agitation 2X.



1. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (1/2).



- 2. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (2/2).
- 34.4 Aspirate.



- 35 Hybridize AMP2.
- 35.1 4-5 drops covering the ample with AMP2.
- 35.2 Incubate for 00:30:00 @ \$ 40 °C .

- 30m

35.3 Wash slides with 1X wash buffer for 2 minutes with slight agitation 2X.

4m

- 1. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (1/2).
- 2. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (2/2).
- 35.4 Aspirate.
- 36 Hybridize AMP3.
- 36.1 4-5 drops covering the sample with AMP3.
- 36.2 Incubate for ♦ 00:15:00 @ \$ 40 °C .





36.3 Wash slides with 1X wash buffer for 2 minutes with slight agitation 2X.



- 1. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (1/2).
- 2. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (2/2).
- 36.4 Aspirate.

Develop Channels (C1-C3)



37

Note

- Don't need to do all 3 channels for each sample just for respective channels, do one channel at a time/slide.
- PC and NC need all 3 Channels.
- Develop 780 channel Last!!

Develop HRP-C2 (570 or 690) signal.

37.1 Add 4-5 drops covering the sample with HRP-C2 (or respective HRP-C#) and incubate for 00:15:00 @ 4 40 °C.

15m

■ Meanwhile.. Dilute Opal dyes → KEEP IN DARK

 \perp 1 μ L Dye (570 or 690) + \perp 1000 μ L TSA buffer

37.2 Wash slides with 1X wash buffer for 2 minutes with slight agitation 2X.

4m

1. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (1/2).

- 2. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (2/2).
- Aspirate.
- 37.3 Add 1st Opal dye and Incubate for \bigcirc 00:30:00 @ \$ 40 °C.

30m

Wash slides with 1X wash buffer for 2 minutes with slight agitation **2X**.

4m

■ Wash slides with 1X wash buffer for 00:02:00 with slight agitation (1/2).

- Wash slides with 1X wash buffer for 00:02:00 with slight agitation (2/2).
- Aspirate.
- 37.5 Add 4-6 drops HRP blocker and incubate for \bigcirc 00:15:00 @ \$ 40 °C .

15m

- 37.6 Wash slides with 1X wash buffer for 2 minutes with slight agitation 2X.

4m

1. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (1/2).

- 2. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (2/2).
- Aspirate.
- 38 Repeat **b** using HRP-C3.
- 39 For 780 Channel
- 39.1 Add HRP-C1 and incubate for 00:15:00 @ \$ 40 °C.

19m

- Wash slides with 1X wash buffer for 2 minutes with slight agitation 2X.
- 1. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (1/2).
- 2. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (2/2).
- Aspirate.
- 39.2 Add ~ 4 120 µL of TSA-DIG per section and incubate for 6 00:30:00 @
- 34m

Room temperature

- Wash slides with 1X wash buffer for 2 minutes with slight agitation 2X.
- 1. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (1/2).
- 2. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (2/2).
- Aspirate.
- 39.3 Add HRP Blocker and incubate for 60 00:15:00 @ 40 °C.



Wash slides with 1X wash buffer for 2 minutes with slight agitation 2X.



- 1. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (1/2).
- 2. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (2/2).
- Aspirate.
- 39.4 Add ~ 4 120 µL of Opal 780 per section and incubate for 6 00:30:00 @

34m

Room temperature



- Wash slides with 1X wash buffer for 2 minutes with slight agitation 2X.
- 1. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (1/2).
- 2. Wash slides with 1X wash buffer for 00:02:00 with slight agitation (2/2).
- Aspirate.



10m

40 Wash w/ PBS 3X.

- Aspirate.
- 41 Apply ~
 ☐ 120 µL of 1X TrueBlack Plus for ○ 00:10:00 @ ☐ Room temperature .



10m

42 Wash w/ PBS 3X.

Aspirate.

DAPI and Mount



- 43 Add 4-5 drops of DAPI to completely cover section.
- 44



30s



- Remove DAPI with absorbent paper \rightarrow tap long side gently to paper.
- 46 Add 3-4 drops of Prolong Gold Mountant.
- 47 Add Coverslip.
- 47.1 Use a forceps to attach coverslip.



- 47.2 Gently press down.
- 47.3 Kim wipe the bottom of the slide and the bench in between samples.
- 48 Store in dark Overnight on absorbent paper (in a drawer).

30s

■ Dry slides in the dark overnight and then Store slides at ▮ 4 °C the next day.



Protocol references

Refer to ACD RNAscope Multiplex Fluorescent v2 Assay for reference (doc #323100).