

# RNAscope® 2.5 HD Duplex Detection Kit (GREEN/RED) Quick Guide

## For FFPE Tissues

#### Introduction

This quick guide is intended for advanced users who are familiar with the procedures in the Formalin-Fixed Paraffin-Embedded (FFPE) Sample Preparation Pretreatment Guide User Manual, Part 1 (Catalog No. 322452-USM) and RNAscope® 2.5 HD Duplex Detection Kit User Manual, Part 2 (Catalog No. 322500-USM). Refer to the user manual for safety guidelines. For every chemical, read the Safety Data Sheet (SDS) and follow handling instructions. Wear appropriate protective eyewear, clothing, and gloves. For the latest services and support information, go to: www.acdbio.com/support.

# Part 1 Prepare and Pretreat Samples

Workflow Steps	
PREPARE FFPE SECTIONS	<ol> <li>Immediately place dissected tissue sample in fresh 10% NBF for 16–32 HRS at ROOM TEMPERATURE (RT).</li> <li>Dehydrate, embed in paraffin, and cut the sample into 5 +/- 1 µm sections. Moun sections on Superfrost® Plus slides.</li> <li>OPTIONAL STOPPING POINT (1). Use sectioned tissue within 3 months. Store sections with desiccants at RT.</li> </ol>
PREPARE SLIDES	Bake Slides
~1.5 HOURS	1. Bake slides in a dry oven for <b>1 HR</b> at <b>60°C</b> .
	OPTIONAL STOPPING POINT (2). Use sectioned tissue within 1 week. Store sections
Bake Slides	with dessicants at RT.
↓ ↓	Deparaffinize FFPE Sections
Deparaffinize FFPE Sections	1. In a fume hood:
	<ul> <li>Fill two Tissue-Tek® Clearing Agent dishes with ~200 mL fresh xylene.</li> </ul>
	<ul> <li>Fill two Tissue-Tek® Staining dishes with ~200 mL fresh 100% EtOH.</li> </ul>
	2. Place slides in a Tissue-Tek® Slide Rack in xylene <b>2 x 5 MIN</b> .
	3. Incubate slides in 100% EtOH <b>2 x 1 MIN</b> .
	4. Remove slides from the rack. Air dry slides for <b>5 MIN</b> at <b>RT</b> .
	OPTIONAL STOPPING POINT (3). Air dry overnight at RT (must use within 24 hrs) or
	proceed directly to the next step.
PRETREAT SAMPLES	Prepare Oven and Reagents (30 MIN at 40°C)
~1-2 HOURS	1. Set $HybEZ^{\mathbb{M}}$ oven to $\mathbf{40^{\circ}C}$ and warm $HybEZ^{\mathbb{M}}$ Humidity Control Tray containing wet
	Humidifying Paper for <b>30 MIN</b> before use. Keep tray warm during the assay.
Prepare Oven and Reagents	2. Prepare 700 mL fresh 1X Target Retrieval in a beaker. Cover with foil, bring to a
↓ A	mild boil, and maintain. Do not boil more than <b>30 MIN</b> before use.
Apply RNAscope® Hydrogen Peroxide	Apply Hydrogen Peroxide (10 MIN at RT)  1. Add ~5-8 drops of Hydrogen Peroxide to each section for 10 MIN at RT.
reroxide	Add ~3-6 diops of Hydrogeth reloxide to each section for <b>10 min</b> at <b>x1</b> .      Place slides into a Tissue-Tek® Slide Rack submerged in distilled water.
↓	<ul><li>3. Wash slides in the distilled water by moving the rack up and down 3–5 times and repeat with fresh distilled water.</li></ul>

Perform Target Retrieval

Create Hydrophobic Barrier

Apply Protease Plus

#### Perform RNAscope® Target Retrieval

- With a pair of forceps very slowly submerge the slide rack into the boiling 1X
   Target Retrieval solution. Refer to Appendix A of the Formalin-Fixed Paraffin-Embedded (FFPE) Sample Preparation Pretreatment Guide User Manual, Part 1 (Cat. No. 322452) for specific pretreatment time, depending on your tissue type.
- 2. *Immediately* transfer hot slide rack to a staining dish containing distilled water.
- 3. Wash slides in the distilled water by moving the rack up and down 3–5 times and repeat with fresh distilled water.
- 4. Wash slides in fresh 100% EtOH by moving the rack up and down 3–5 times and air dry.

#### **Create Hydrophobic Barrier**

Draw 2–4 times around tissue using the Immedge<sup>™</sup> hydrophobic barrier pen. Dry completely ~2 MIN or OVERNIGHT at RT.

#### **Apply Protease Plus**

- Place slides in the HybEZ<sup>™</sup> Slide Rack, and add ~5 drops of Protease Plus to each section.
- Place the HybEZ<sup>™</sup> Slide Rack in the prewarmed HybEZ<sup>™</sup> Humidity Control Tray.
   Seal tray and insert back into the HybEZ<sup>™</sup> Oven. Incubate at 40°C for 30 MIN.

**Note:** If needed, prepare RNAscope® 2.5 assay materials during this step.

3. Wash slides in the distilled water by moving the rack up and down 3–5 times and repeat with fresh distilled water.

Part 2: RNAscope® 2.5 Duplex Assay

#### Workflow Steps 1. Prepare 3 L of 1X Wash Buffer by adding 2.94 L distilled water and 1 bottle (60 PREPARE THE MATERIALS mL) of 50X Wash Buffer to a large carboy. Mix well. Warm 50X Wash Buffer up to ~10-30 MIN 40°C for 10-20 MIN before making 1X Wash Buffer. 2. Prepare 50% Hematoxylin and 0.02% Ammonia water. 3. Equilibrate remove Amps 1-10 from the refrigerator and keep at RT**Prepare Probes** 4. Warm probes for 10 MIN at 40°C, then cool to RT. 5. Briefly spin the C2 probe. 6. Mix 1:50 ratio of C2 probe to C1 probe by pipetting 1 volume of C2 probe to 50 volumes of C1 probe into a tube. Invert the tube several times. 7. Mixed probes can be stored at 4°C for up to 6 months. **RUN THE ASSAY** Hybridize Probe (2 HRS at 40°C) ~7 Hours 1. Remove excess liquid from slides, place in the HybEZ<sup>™</sup> Slide Rack, and add ~4 drops probe to each section. 2. Insert the sealed tray containing $HybEZ^{™}$ Slide Rack back into the $HybEZ^{™}$ Oven for Hybridize Probe 2 HRS at 40°C. Remove slide rack. Hybridize Amp 1 3. Wash slides in 1X Wash Buffer for **2 MIN** at **RT**. Repeat with fresh 1X Wash Buffer. Hybridize Amp 1 (30 MIN at 40°C) Hybridize Amp 2 1. Remove excess liquid from slides, place in the HybEZ<sup>™</sup> Slide Rack, and add ~4 drops Amp 1 to each section. Hybridize Amp 3 Insert the sealed tray containing HybEZ<sup>™</sup> Slide Rack into the HybEZ<sup>™</sup> Oven for 30 MIN at 40°C. Remove slide rack. Hybridize Amp 4 Wash slides in 1X Wash Buffer for 2 MIN at RT. Repeat with fresh 1X Wash Buffer.

Hybridize Amp 5

Hybridize Amp 6

Hybridize Amp 6

Detect the Red Signal

Hybridize Amp 7

Hybridize Amp 8

Hybridize Amp 9

Hybridize Amp 10

Detect the Green Signal

Counterstain the Slides

Mount the Slides

#### Hybridize Amp 2 (15 MIN at 40°C)

- Remove excess liquid from slides, place in the HybEZ<sup>™</sup> Slide Rack, and add ~4 drops Amp 2 to each section.
- Insert the sealed tray containing HybEZ<sup>™</sup> Slide Rack into the HybEZ<sup>™</sup> Oven for 15 MIN at 40°C. Remove slide rack.
- 3. Wash slides in 1X Wash Buffer for **2 MIN** at **RT**. Repeat with fresh buffer.

#### Hybridize Amp 3 (30 MIN at 40°C)

- Remove excess liquid from slides, place in the HybEZ<sup>™</sup> Slide Rack, and add ~4 drops Amp 3 to each section.
- Insert the sealed tray containing HybEZ<sup>™</sup> Slide Rack into the HybEZ<sup>™</sup> Oven for 30 MIN at 40°C. Remove slide rack.
- 3. Wash slides in 1X Wash Buffer for **2 MIN** at **RT**. Repeat with fresh buffer.

### Hybridize Amp 4 (15 MIN at 40°C)

- Remove excess liquid from slides, place in the HybEZ<sup>™</sup> Slide Rack, and add ~4 drops Amp 4 to each section.
- 2. Insert the sealed tray containing HybEZ<sup>™</sup> Slide Rack into the HybEZ<sup>™</sup> Oven for **15 MIN** at **40°C**. Remove slide rack, but do *not* place the tray back into the oven.
- 3. Wash slides in 1X Wash Buffer for **2 MIN** at **RT**. Repeat with fresh buffer.

### Hybridize Amp 5 (30 MIN at RT)

- Remove excess liquid from slides, place in the HybEZ<sup>™</sup> Slide Rack, and add ~4 drops Amp 5 to each section.
- 2. Incubate the sealed tray containing HybEZ™ Slide Rack for **30 MIN** at **RT**.
- 3. Wash slides in 1X Wash Buffer for **2 MIN** at **RT**. Repeat with fresh buffer.

#### Hybridize Amp 6 (15 MIN at RT)

- Remove excess liquid from slides, place in the HybEZ<sup>™</sup> Slide Rack, and add ~4 drops Amp 6 to each section.
- 2. Incubate the sealed tray containing HybEZ<sup>™</sup> Slide Rack for **15 MIN** at **RT**.
- 3. Wash slides in 1X Wash Buffer for **2 MIN** at **RT**. Repeat with fresh buffer.

#### Detect the Signal (10 MIN at RT)

- 1. Briefly spin RED-B and mix 1 volume of RED-B to 60 volumes of RED-A (must use within **3–5 MIN**). For examples add 2.5 μL of Red-B to 150 μL of Red-A per section.
- 2. Remove excess liquid from slides, place in the HybEZ<sup>™</sup> Slide Rack, and pipette ~120 µL of RED solution onto each tissue section.
- 3. Incubate sealed tray containing HybEZ™ Slide Rack for 10 MIN at RT.
- 4. Wash slides in 1X Wash Buffer for **2 MIN** at **RT**. Repeat with fresh buffer.

#### Hybridize Amp 7 (15 MIN at 40°C)

- Remove excess liquid from slides, place in the HybEZ<sup>™</sup> Slide Rack, and add ~4 drops Amp 7 to each section.
- 2. Incubate the sealed tray containing HybEZ™ Slide Rack for 15 MIN at 40°C.
- 3. Wash slides in 1X Wash Buffer for **2 MIN** at **RT**. Repeat with fresh buffer.

#### Hybridize Amp 8 (30 MIN at 40°C)

- Remove excess liquid from slides, place in the HybEZ<sup>™</sup> Slide Rack, and add ~4 drops Amp 8 to each section.
- Incubate the sealed tray containing HybEZ<sup>™</sup> Slide Rack for 30 MIN at 40°C.
- 3. Wash slides in 1X Wash Buffer for **2 MIN** at **RT**. Repeat with fresh buffer.

#### Hybridize Amp 9 (30 MIN at RT)

- Remove excess liquid from slides, place in the HybEZ<sup>™</sup> Slide Rack, and add ~4 drops Amp 9 to each section.
- 2. Incubate the sealed tray containing HybEZ<sup>™</sup> Slide Rack for **30 MIN** at **RT**.

	3. Wash slides in 1X Wash Buffer for <b>2 MIN</b> at <b>RT</b> . Repeat with fresh buffer.
	Hybridize Amp 10 (15 MIN at RT)
	<ol> <li>Remove excess liquid from slides, place in the HybEZ<sup>™</sup> Slide Rack, and add ~4</li> </ol>
	drops Amp 10 to each section.
	<ol> <li>Incubate the sealed tray containing HybEZ™ Slide Rack for 15 MIN at RT.</li> </ol>
	3. Wash slides in 1X Wash Buffer for <b>2 MIN</b> at <b>RT</b> . Repeat with fresh buffer.
	Detect the Signal (10 MIN at RT)
	1. Briefly spin GREEN-B and mix 1 volume of Green-B to 50 volumes of GREEN-A
	(must use within <b>3–5 MIN</b> ). For examples add 3 $\mu L$ of GREEN-B to 150 $\mu L$ of
	GREEN-A per section.
	2. Remove excess liquid from slides, place in the HybEZ™ Slide Rack, and pipette
	~120 µL of GREEN solution onto each tissue section.
	3. Incubate sealed tray containing $HybEZ^{T}$ Slide Rack for 10 MIN at RT.
	4. Remove solution from slides and wash <b>5 MIN</b> with 1X wash buffer
	5. Rinse quickly with distilled water.
	Counterstain the Slides (30 Sec at RT)
	1. Place slides in 50% Hematoxylin I for <b>30 Sec</b> at <b>RT</b> . Wash 3–5 times in tap water
	and repeat with fresh tap water.
	2. Wash slides 10 SEC in 0.02% Ammonia water, and then wash 3–5 times in tap
	water.
	Mount the Slides
	1. Dry slides in a <b>60°C</b> dry oven for <b>15 MIN</b> .
	2. Dip the slides into FRESH pure xylene and immediately place 1–2 drops of
	EcoMount on the slide before the xylene dries. Place a coverslip over the section.
	3. Air dry for <b>30 MIN</b> .
EVALUATE THE RESULTS	Examine tissue sections under a standard bright field microscope at 20–40X magnification

# **Troubleshooting**

For troubleshooting information, please contact technical support at support.acd@bio-techne.com.

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