

HCRTM RNA-FISH (v3.0)/IF protocol for sample on slide

This protocol has not been validated for all sample types and should only be used as a template.

Technical Support support@molecularinstruments.com

Safety Data Sheets (SDS) www.molecularinstruments.com/safety-v3

Patents

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Order one HCRTM IF kit per target protein Order one HCRTM **RNA-FISH** (v3.0) kit per target **RNA**

Example 2-Plex Experiment

- HCRTM RNA-FISH (v3.0) kit for target mRNA1
 - $\circ~$ HCRTM Probe (v3.0): target mRNA1 for use with amplifier B1
 - HCR[™] Amplifier (v3.0): B1-647
 - O HCR[™] RNA-FISH Buffers (v3.0): HCR[™] Probe Hybridization Buffer (v3.0), HCR[™] Probe Wash Buffer (v3.0), HCR[™] Amplifier Buffer (v3.0)
- HCRTM IF kit for target Protein1
 - 1° Ab: Mouse Anti-Protein1 (your own 1° antibody)
 - HCRTM 2° Antibody Probe: Donkey Anti-Mouse for use with amplifier B2
 - HCR™ Amplifier (v3.0): B2-488
 - HCRTM IF Buffers: HCRTM Antibody Buffer, HCRTM Amplifier Buffer (v3.0)

Storage conditions

- Store HCRTM Probes (v3.0), HCRTM 2° Antibody Probes, HCRTM Amplifiers (v3.0), HCRTM Probe Hybridization Buffer (v3.0), and HCRTM Probe Wash Buffer (v3.0) at -20 °C.
- Store HCRTM Antibody Buffer and HCRTM Amplifier Buffer (v3.0) at 4 °C.
- On the bench top, keep stock solutions on ice.
- Make sure all solutions are well mixed before use.



Sample preparation

Samples should be prepared in the same manner as for a traditional immunohistochemistry, up to the primary antibody application step. This may include permeabilization and antigen retrieval. Then proceed with the protocol described below.

Multiplexed HCRTM IF + HCRTM RNA-FISH (v3.0) protocol

Protein detection stage

- 1. Block tissue by applying 200 μ L of HCRTM Antibody Buffer on top of the sample. Incubate at room temperature for 1 h in a humidified chamber.
- 2. Prepare working concentration of primary antibodies in HCR[™] Antibody Buffer. Prepare 100 μL per section. NOTE: *follow manufacturer's guidelines for primary antibody working concentration.*
- 3. Drain slide by blotting edges on a Kimwipe and wipe around the section with another Kimwipe.
- 4. Add primary antibody solution to each section and incubate overnight (>12 h) at 4 °C in a humidified chamber. NOTE: *Incubation may be optimized (e.g., 1–2 h at room temperature) depending on sample type and thickness.*
- 5. Remove excess antibodies by immersing slide in $1 \times PBST$ at room temperature for 3×5 min.
- 6. Prepare 1 µg/mL working concentration of HCR[™] 2° Antibody Probes in HCR[™] Antibody Buffer. Prepare 100 µL per section.
 NOTE: Concentration may be optimized depending on protein target and primary antibody.
- 7. Drain slide by blotting edges on a Kimwipe and wipe around the section with another Kimwipe.
- 8. Add secondary antibody solution to each section and incubate for 1 h at room temperature in a humidified chamber.
- 9. Remove excess antibodies by immersing slide in $1 \times PBST$ at room temperature for 3×5 min.
- 10. Proceed to **RNA detection stage** for co-detection of protein and RNA.



RNA detection stage

- 1. Drain slide by blotting edges on a Kimwipe and wipe around the section with another Kimwipe.
- 2. Post-fix sample by adding 200 μ L of 4% formaldehyde on the tissue. CAUTION: *use formaldehyde with extreme care as it is a hazardous material.*
- 3. Incubate slides for 10 min at room temperature.
- 4. Immerse slides for 2×5 min in PBST.
- 5. Immerse slides for 5 min in $5 \times$ SSCT.
- 6. Pre-warm a humidified chamber to 37 °C.
- 7. Drain slide by blotting edges on a Kimwipe and wipe around the section with another Kimwipe.
- 8. Add 200 μL of HCRTM Probe Hybridization Buffer (v3.0) on top of the tissue sample.
 CAUTION: HCRTM Probe Hybridization Buffer (v3.0) contains formamide, a hazardous material.
 NOTE: pre-heat HCRTM Probe Hybridization Buffer (v3.0) to 37 °C before use.
- 9. Pre-hybridize for 10 min inside the humidified chamber.
- 10. Prepare a 16 nM probe solution by adding 1.6 pmol of each HCRTM Probe (v3.0) (e.g. 1.6 μL of 1 μM stock) to 100 μL of HCRTM Probe Hybridization Buffer (v3.0) at 37 °C.
 NOTE: This is the amount of probe needed for each target on a single slide using 100 μL of incubation volume.
- 11. Remove the pre-hybridization solution and drain excess buffer on slide by blotting edges on a Kimwipe.
- 12. Add 100 μ L of the probe solution on top of the tissue sample.
- 13. Place a coverslip on the sample and incubate overnight (>12 h) in the 37 °C humidified chamber.
- 14. Immerse slide in HCRTM Probe Wash Buffer (v3.0) at 37 °C to float off coverslip. CAUTION: *HCRTM Probe Wash Buffer (v3.0) contains formamide, a hazardous material.*
- 15. Remove excess probes by incubating slide at 37 °C in:
 - (a) 75% of HCRTM Probe Wash Buffer (v3.0) / 25% $5 \times$ SSCT for 15 min
 - (b) 50% of HCRTM Probe Wash Buffer (v3.0) / 50% $5 \times$ SSCT for 15 min
 - (c) 25% of HCRTM Probe Wash Buffer (v3.0) / 75% $5 \times$ SSCT for 15 min
 - (d) 100% $5 \times$ SSCT for 15 min

NOTE: Wash solutions should be pre-heated to 37 °C before use.

16. Proceed to Amplification stage.



Amplification stage

- 1. Immerse slide in $5 \times$ SSCT at room temperature for 5 min.
- 2. Drain slide by blotting edges on a Kimwipe and wipe around the section with another Kimwipe.
- Add 200 µL of HCR[™] Amplifier Buffer (v3.0) on top of the tissue sample and pre-amplify in a humidified chamber for 30 min at room temperature.
 NOTE: equilibrate HCR[™] Amplifier Buffer (v3.0) to room temperature before use.
- 4. Separately prepare 6 pmol of hairpin h1 and 6 pmol of hairpin h2 by snap cooling 2 μL of 3 μM stock (heat at 95 °C for 90 seconds and cool to room temperature in a dark drawer for 30 min). NOTE: Hairpins h1 and h2 are provided in hairpin storage buffer ready for snap cooling. h1 and h2 should be snap cooled in separate tubes. This is the amount of hairpins needed for each target on a single slide using 100 μL of incubation volume.
- 5. Prepare a 60 nM hairpin solution by adding all snap-cooled h1 hairpins and snap-cooled h2 hairpins to 100 μ L of HCRTM Amplifier Buffer (v3.0) at room temperature per section.
- 6. Remove the pre-amplification solution and drain excess buffer on slide by blotting edges on a Kimwipe.
- 7. Add 100 μ L of the hairpin solution on top of the tissue sample.
- 8. Incubate overnight (>12 h) in a dark humidified chamber at room temperature.
- 9. Remove excess hairpins by immersing slide in $5 \times$ SSCT at room temperature for:
 - (a) $2 \times 5 \min$
 - (b) $2 \times 15 \text{ min}$
 - (c) $1 \times 5 \min$
- 10. Drain slide by blotting edges on a Kimwipe and dry around the section with another Kimwipe.
- 11. Add 50–100 μ L of antifade mounting reagent on top of the sample.
- 12. Place a coverslip on top for microscopy.



Buffer recipe

PBST

1× PBS 0.1% Tween 20

 $\frac{5 \times \text{SSCT}}{5 \times \text{ sodium chloride sodium citrate (SSC)}}$ 0.1% Tween 20 $\frac{\text{For 50 mL of solution}}{5 \text{ mL of } 10 \times \text{PBS}}$ $500 \ \mu\text{L of } 10\% \text{ Tween } 20$ Fill up to 50 mL with ultrapure H₂O

NOTE: avoid using calcium chloride and magnesium chloride in PBS as this leads to increased autofluorescence in the samples.



S1 HCRTM Technology Citation Notes

For citation, please select from the list below as appropriate for your application:

• HCRTM RNA-ISH

HCR[™] RNA in situ hybridization (RNA-ISH) offers unmatched performance, robustness, and versatility imaging RNA targets in diverse organisms and sample types (Choi et al., 2010, Choi et al., 2014, Choi et al., 2018):

- HCRTM RNA-FISH

HCR[™] RNA-FISH enables 10-plex, quantitative, high-resolution RNA fluorescence in situ hybridization (RNA-FISH) with automatic background suppression throughout the protocol for dramatically enhanced performance (signal-to-background, quantitative precision, single-molecule fidelity) and ease-of-use (no probe set optimization for new targets and organisms).

- Enzymatic HCRTM RNA-CISH/RNA-FISH

Enzymatic HCRTM RNA-ISH integrates enzymatic signal amplification to enable extreme-sensitivity RNA imaging using either chromogenic or fluorescent staining (RNA-CISH/RNA-FISH). In tissue sections, entirely protease-free workflows preserve sample morphology and maintain protein target integrity, enabling seamless compatibility with existing immunohistochemistry (IHC)/immunofluorescence (IF) assays. HCRTM RNA-CISH offers the convenience of brightfield microscopy and the option of archival staining.

• 10-Plex HCRTM Spectral Imaging

HCR[™] RNA-FISH/IF enables quantitative high-resolution imaging of 10 RNA and/or protein targets with 1step HCR[™] signal amplification for all targets simultaneously. The method is suitable even for whole-mounts and delicate samples as it requires no repeated staining, imaging, registration, or stripping (Schulte et al., 2024).

• HCRTM RNA-FISH/IF

HCR[™] RNA-FISH/IF enables a unified approach to multiplex, quantitative, high-resolution RNA fluorescence in situ hybridization (RNA-FISH) and protein immunofluorescence (IF), with quantitative 1-step enzymefree signal amplification performed for all RNA and protein targets simultaneously (Schwarzkopf et al., 2021).

• **HCR**TM **IF**

HCR[™] IF enables multiplex, quantitative, high-resolution protein immunofluorescence (IF) in highly autofluorescent samples (e.g., FFPE brain tissue sections) (Schwarzkopf et al., 2021).

• Subcellular Quantitative RNA and Protein Imaging

HCR[™] RNA-FISH enables analog relative quantitation of RNA and/or protein targets with subcellular resolution in the anatomical context of thick autofluorescent samples (e.g., whole-mount vertebrate embryos) (Trivedi et al., 2018, Choi et al., 2018, Schwarzkopf et al., 2021).

• Single-Molecule Quantitative RNA Imaging

HCR[™] RNA-FISH enables digital RNA absolute quantitation with single-molecule resolution in the anatomical context of thick autofluorescent samples (e.g., 0.5 mm adult mouse brain sections) (Shah et al., 2016, Choi et al., 2018).

Read-Out/Read-In Analysis Framework

The read-out/read-in analysis framework enables bidirectional quantitative discovery in an anatomical context (Trivedi et al., 2018).

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Protocols in Diverse Sample Types

Protocols for HCR[™] RNA-FISH and/or IF in diverse sample types are adapted from the zoo paper (Choi et al., 2016):

- bacteria in suspension
- FFPE human tissue sections
- generic sample in solution
- generic sample on a slide
- mammalian cells on a slide
- mammalian cells in suspension
- whole-mount chicken embryos
- whole-mount fruit fly embryos
- whole-mount mouse embryos
- \circ whole-mount nematode larvae
- whole-mount sea urchin embryos
- whole-mount zebrafish embryos and larvae

• HCRTM RNA Flow Cytometry

HCR[™] RNA Flow Cytometry enables analog RNA relative quantitation for high-throughput expression profiling of mammalian cells and bacteria without the need to engineer reporter lines (Choi et al., 2018).

• HCRTM Northern Blots

HCR[™] Northern Blots enable simultaneous quantification of RNA target size and abundance with automatic background suppression throughout the protocol (Schwarzkopf & Pierce, 2016).

• HCRTM Amplifiers

HCR[™] Amplifiers enable multiplex, quantitative, 1-step, isothermal, enzyme-free signal amplification in diverse technological settings (Dirks & Pierce, 2004).