

HCR™ northern blot protocol

Technical support

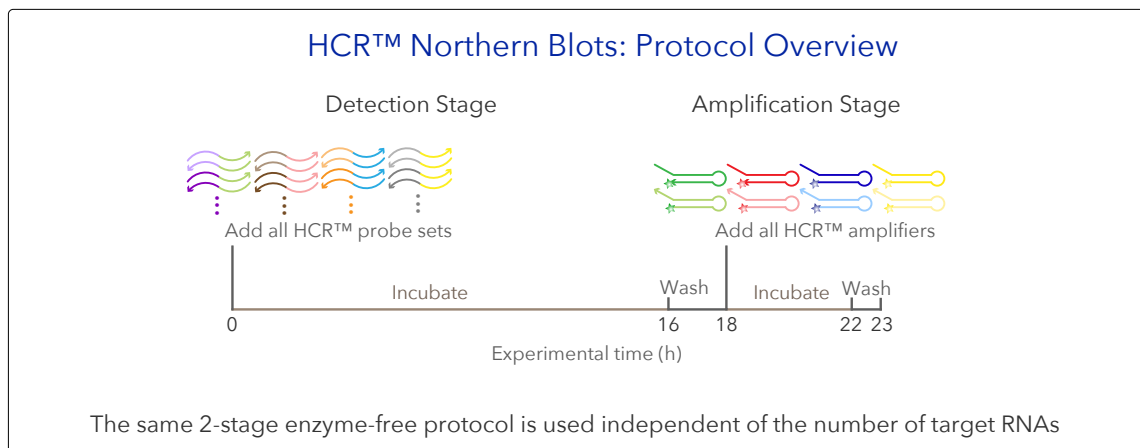
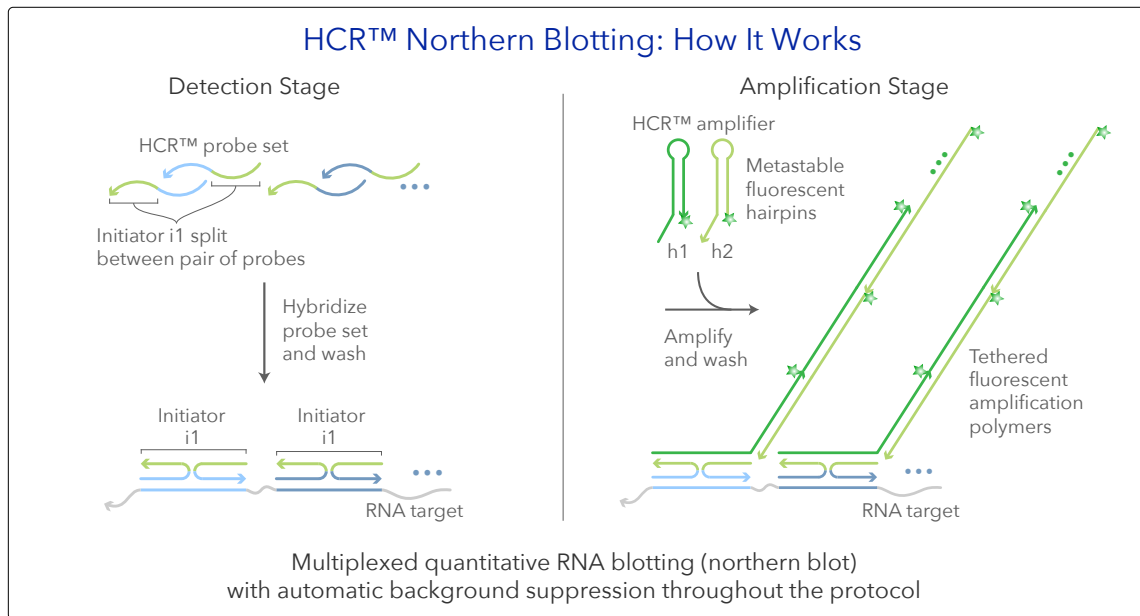
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HCR™ Northern Blots

Multiplexed quantitative RNA blotting

Multiplexed Experiment

- Order one HCR™ RNA-FISH bundle per target RNA

Example 2-Plex Experiment

- HCR™ RNA-FISH bundle for target mRNA1
 - HCR™ probe set: target mRNA1 for use with amplifier B1
 - HCR™ amplifier: B1-647
 - HCR™ RNA-FISH buffers: probe hybridization buffer, probe wash buffer, amplification buffer (for use with all bundles)
- HCR™ RNA-FISH bundle for target mRNA2
 - HCR™ probe set: target mRNA2 for use with amplifier B2
 - HCR™ amplifier: B2-488

Storage conditions

- Store HCR™ probe sets, HCR™ amplifiers, HCR™ RNA-FISH probe hybridization buffer, and HCR™ RNA-FISH probe wash buffer at -20 °C.
- Store HCR™ amplification buffer at 4 °C.
- On the bench top, keep stock solutions on ice.
- Make sure all solutions are well mixed before use.

Denaturing agarose gel electrophoresis

1. Prepare a 1% denaturing agarose gel.
NOTE: The gel should be ≤ 6 mm for efficient RNA transfer.
NOTE: We recommend using NorthernMax™ denaturing gel buffer for preparing denaturing agarose gels.
2. Mix RNA samples in a 1:1 ratio with formamide and heat to 65 °C for 15 min prior to gel loading.
CAUTION: formamide is a hazardous material.
3. Run RNA samples (typically 1–20 μ g) at 5V per cm in 1 \times NorthernMax™ running buffer until your targets of interest are well-resolved.
NOTE: The distance measured is the distance between the electrodes in the electrophoresis chamber.
NOTE: For targets with low or unknown expression levels we recommend starting with 10–20 μ g of total RNA.

RNA transfer and crosslinking

1. Wash the gel 4 \times 5 min, in ultrapure H₂O.
2. Wet membrane in ultrapure H₂O and equilibrate for 5 min in 20 \times SSC.
NOTE: A positively charged nylon membrane is recommended.
3. Perform a capillary transfer of RNA onto the membrane in 20 \times SSC.
NOTE: We recommend using the TurboBlotter Rapid Downward Transfer System. Alternative transfer methods may be used.
4. Wash the membrane for 5 min in 2 \times SSC.
5. Crosslink the RNA to the membrane by incubating at 80 °C for 2 h.
NOTE: Alternative RNA crosslinking methods may be used.

Multiplexed HCR™ northern blot protocol

Detection stage

1. Let V denote the volume of buffer to be used for blots, calculated as:

- (a) 1 mL per 10 cm² of membrane
- (b) rounded up to the nearest 0.5 mL
- (c) no less than 1.5 mL

For example, $V = 3.5$ mL for a membrane that is 32 cm² and $V = 1.5$ mL for a membrane that is 9 cm².

2. Pre-heat probe hybridization buffer to 37 °C.

CAUTION: *Probe hybridization buffer contains formamide, a hazardous material.*

3. Pre-hybridize blot in ($V - 0.5$ mL) of probe hybridization buffer in a hybridization bottle for 30–60 min at 37 °C in a rolling hybridization incubator.

4. Prepare probe solution by adding each probe set into a total volume of 0.5 mL of probe hybridization buffer at 37 °C such that each probe set will be at a final concentration of 5 nM in volume V .

5. Add the probe solution to the pre-hybridization solution and incubate blot overnight at 37 °C in a rolling hybridization incubator.

6. Remove excess probes by washing four times using pre-heated 37 °C wash solutions (volume $2V$ per wash):

- (a) two low-stringency washes ($2 \times$ SSC, 0.1% SDS; 5 min at 37 °C)
- (b) two high-stringency washes ($0.2 \times$ SSC, 0.1% SDS; 15 min at 37 °C)

Amplification stage

1. Pre-heat amplification buffer to 37 °C.

2. Pre-amplify blot in ($V - 0.5$ mL) of amplification buffer in a hybridization bottle for 30–60 min at 37 °C in a rolling hybridization incubator.

3. Separately prepare hairpin h1 and hairpin h2 by snap cooling (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. Snap cool an amount of each hairpin corresponding to 30 nM final concentration in a final volume V . h1 and h2 should be snap cooled in separate tubes.*

4. Prepare hairpin solution by adding all snap-cooled hairpins into a final volume of 0.5 mL amplification buffer at 37 °C.

5. Add the hairpin solution to the pre-amplification solution and incubate blot for 4 h at 37 °C in the dark in a rolling hybridization incubator.

NOTE: *Amplification time can be increased beyond 4 h if desired.*

NOTE: *Amplification may alternatively be carried out at room temperature.*

6. Remove excess hairpins by washing in pre-heated $5 \times$ SSCT (volume $2V$ per wash): 2×15 min at 37 °C in the dark.

NOTE: *If amplification is performed at room temperature, hairpin washes should also be performed at room temperature.*

Buffer recipes for sample preparation

10% SDS

10% SDS

For 100 mL of solution

10 g of SDS powder

Fill up to 100 mL with ultrapure H₂O

Heat solution to 68 °C to dissolve powder if necessary.

2× SSC, 0.1% SDS

2× sodium chloride sodium citrate (SSC)

0.1% SDS

For 100 mL of solution

10 mL of 20× SSC

1 mL of 10% SDS

Fill up to 100 mL with ultrapure H₂O

0.2× SSC, 0.1% SDS

0.2× sodium chloride sodium citrate (SSC)

0.1% SDS

For 100 mL of solution

1 mL of 20× SSC

1 mL of 10% SDS

Fill up to 100 mL with ultrapure H₂O

5× SSCT

5× sodium chloride sodium citrate (SSC)

0.1% Tween 20

For 40 mL of solution

10 mL of 20× SSC

400 μL of 10% Tween 20

Fill up to 40 mL with ultrapure H₂O

Reagents and supplies

NorthernMax™ denaturing gel buffer (10X) (ThermoFisher Scientific, Cat. # AM8676)
NorthernMax™ (10X) running buffer (ThermoFisher Scientific, Cat. # AM8671)
Nylon membrane, positively-charged (Roche, Cat. # 11209272001)
3 mm Whatman chromatography paper (GE Healthcare Life Sciences, Cat. # 3030-6188)
Whatman TurboBlotter (Fisher Scientific, Cat. # 09-301-184)
Whatman gel blotting paper (GE Healthcare Life Sciences, Cat. # 10427805)
Whatman GB003 filter paper (GE Healthcare Life Sciences, Cat. # 10427812)
Hybridization bottle (Wheaton, Cat. # 805000 or 805021)
Rolling hybridization incubator (SciGene, Cat. # 1040-50-1)

HCR™ Technology Citation Notes

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

- **HCR™ IF + HCR™ RNA-FISH**

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

- **HCR™ IF**

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

- **HCR™ RNA-FISH (v3.0)**

Third-generation [HCR™ RNA-FISH \(v3.0\)](#) enables multiplexed, quantitative, high-resolution RNA fluorescence in situ hybridization (RNA-FISH) with [automatic background suppression throughout the protocol](#) for dramatically enhanced performance (signal-to-background, subcellular quantitative RNA imaging precision, single-molecule quantitative RNA imaging fidelity) and ease-of-use (no probe set optimization for new targets and organisms) ([Choi et al., 2018](#)). Quantitative analysis modes:

- [Subcellular quantitative RNA imaging](#): analog mRNA relative quantitation with subcellular resolution in the anatomical context of thick autofluorescent samples.
- [Single-molecule quantitative RNA imaging](#): digital mRNA absolute quantitation with single-molecule resolution in the anatomical context of thick autofluorescent samples.
- [Quantitative RNA flow cytometry](#): analog mRNA relative quantitation for high-throughput expression profiling of mammalian cells and bacteria.

[Protocols for HCR™ RNA-FISH \(v3.0\)](#) in diverse organisms are adapted from the Zoo paper.

- **Subcellular quantitative RNA imaging**

[Subcellular quantitative RNA imaging](#) enables mRNA relative quantitation with subcellular resolution in the anatomical context of thick autofluorescent samples (e.g., whole-mount vertebrate embryos). The [read-out/read-in analysis framework](#) enables bidirectional quantitative discovery in an anatomical context ([Trivedi et al., 2018](#)).

- **Zoo paper**

Protocols for multiplexed mRNA imaging in diverse sample types ([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
- whole-mount nematode larvae
- whole-mount sea urchin embryos
- whole-mount zebrafish embryos and larvae

- **Single-molecule quantitative RNA imaging**

[Single-molecule quantitative RNA imaging](#) enables 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](#)).

- **HCR™ northern blots**

[HCR™ northern blots](#) enable multiplexed quantification of RNA target size and abundance for up to 5 target RNAs ([Schwarzkopf & Pierce, 2016](#)).

- **HCR™ RNA-FISH (v2.0)**

Second-generation in situ HCR™ RNA-FISH technology (v2.0) using DNA HCR™ probes and DNA HCR™ amplifiers: 10× increase in signal, 10× reduction in cost, dramatic increase in reagent durability ([Choi et al., 2014](#)).

- **HCR™ RNA-FISH (v1.0)**

First-generation HCR™ RNA-FISH technology (v1.0) using RNA HCR™ probes and RNA HCR™ amplifiers: multiplexed mRNA imaging in whole-mount vertebrate embryos with simultaneous signal amplification for up to 5 target mRNAs ([Choi et al., 2010](#)).

- **HCR™ technology**

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