

# FISH Probes – Automated Hybridization Protocol

## Notes

- Protocol can be used with all FISH probes – controls, gene specific, custom FISH probes.
- Solutions can be made prior to the procedure.
- Further optimization of the protocol may be required.

## Required Reagents & Equipment (Not Supplied)

HYBrite or ThermoBrite

Absorbent material

dH<sub>2</sub>O

Wash Solution 1 (WS1) – 0.3% Igepal (Sigma CA-630) or NP-40 / 0.4 x SSC

Wash Solution 2 (WS2) – 0.1% Igepal (Sigma CA-630) or NP-40 / 2 x SSC

DAPI with Antifade

## Automated HYBrite / ThermoBrite Protocol

1. Turn on HYBrite / ThermoBrite.
2. Set program. Please see guide below.
3. Presoak absorbent material in dH<sub>2</sub>O and position in HYBrite / ThermoBrite.
4. Add 10 µl probe mixture to slide (2 µl probe + 8 µl hybridization buffer, or for multiple individually supplied probes per slide use 2 µl of each probe plus hybridization buffer for a total volume of 10 µl per slide).
5. Apply clean 22 mm<sup>2</sup> coverslip to slide.
6. Apply rubber cement on edges of coverslip to seal.
7. Place in HYBrite or ThermoBrite and close lid.
8. Start program. Hybridization will take at least 16 hours. See below for program recommendations.
9. Pre-warm WS1 (0.3% Igepal (Sigma CA-630) or NP-40 / 0.4 x SSC) to 73°C.
10. Remove coverslip. Place in WS1, agitating for approximately 10 seconds then let stand for exactly 2 minutes.
11. Transfer to WS2 (0.1% Igepal (Sigma CA-630) or NP-40 / 2 x SSC) at room temperature for 1 minute.
12. Let dry in dark.
13. Apply 10 µl DAPI with Antifade and 22 mm<sup>2</sup> coverslip.
14. Wait 15-30 minutes then visualize under microscope using the appropriate filter sets.

## HYBrite / ThermoBrite Program Guide

### *Peripheral Blood Preparations*

Denature at 72-73°C for 2 minutes. Hybridize at 37°C for at least 16 hours.

### *Paraffin Embedded Tissue Sections (after pretreatment)*

Denature at 83°C for 3 minutes. Hybridize at 37°C for at least 16 hours. May require troubleshooting.

## Recommendations

### *Optional FISH Pretreatment*

We recommend the Abbott Molecular FISH Pretreatment Reagent Kit.

### *Paraffin Pretreatment*

Please see the Empire Genomics Paraffin Embedded Tissue Sample Slide Processing Protocol.

Alternatively we recommend the Abbott Molecular Paraffin Pretreatment Reagent Kits: I, II or III.

## References

Barch MJ, Knutsen T, Spurbeck JL. The AGT Cytogenetics Laboratory Manual, Third Edition. Lippincott-Raven Philadelphia. 1991.



Empire Genomics  
700 Michigan Avenue, Suite 200  
Buffalo, NY 14203  
1-800-715-5880  
info@empiregenomics.com  
www.empiregenomics.com

### Dye Specification Sheet

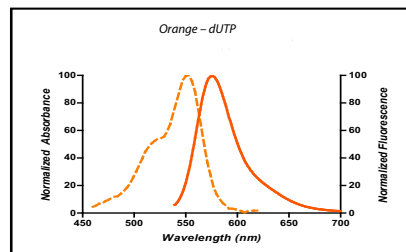
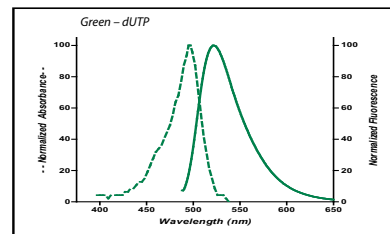
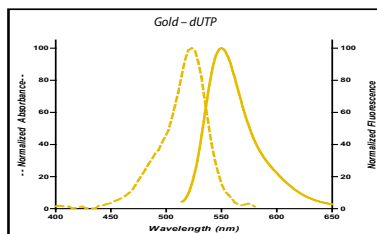
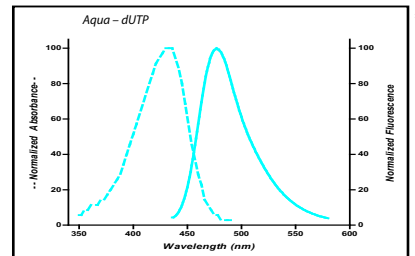
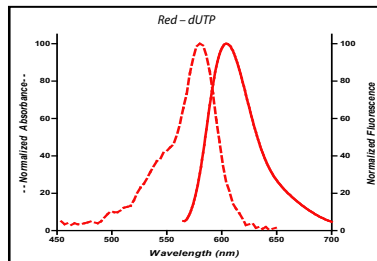
|                    |                                |               |             |                     |           |
|--------------------|--------------------------------|---------------|-------------|---------------------|-----------|
| BAC DNA Library    | RP11®, RP23®                   |               |             |                     |           |
| Cot-1 DNA          | 15 µg (3 µg/reaction)          |               |             |                     |           |
| Diluent            | 1x Hyb Buffer                  |               |             |                     |           |
| Label              | Red-dUTP                       | Green-dUTP    | Orange-dUTP | Gold-dUTP           | Aqua-dUTP |
| Fluorophore        | 5-ROX (5-Carboxyl-x-rhodamine) | 5-Fluorescein | 5-TAMRA     | Carboxyrhodamine 6G | Aqua      |
| Color              | Red                            | Green         | Orange      | Gold                | Aqua      |
| Absorbance Maximum | 580 nm                         | 491 nm        | 548 nm      | 525 nm              | 418 nm    |
| Emission Maximum   | 599 nm                         | 515 nm        | 573 nm      | 551 nm              | 467 nm    |

#### INSTRUCTIONS

- Gently vortex and centrifuge tube prior to use
- Store at -20°C in a manual defrost freezer
- Protect from light
- Minimize freeze-thaw cycles

#### FISH PROTOCOL

Please refer to the FISH & Hybridization Protocol on our website:  
<http://www.empiregenomics.com/docs/HybridizationQuickReference-EmpireGenomics.pdf>



\* Please Note: The human eye visualizes the Aqua wavelength more poorly than other regions of the visible light spectrum (as above). Consequently, when choosing to use an aqua probe, it is best to use it with a target that hybridizes strongly. For example, in our own experiences we have had better success with centromere probes compared to locus probes. Our Aqua probes have been benchmarked against the leading competitors and we are as bright as or brighter than they are. This material has passed our Quality Control processes and meets performance benchmarks. We offer a variety of colors for FISH probe labeling and if you want a probe with a stronger signal we would suggest you consider using green, gold, orange or red ones. We cannot guarantee the performance you will experience with the aqua dye as a result of the many variables which can affect its performance

#### For Research Use Only

This product is sold by Empire Genomics LLC for research purposes only and is not intended for diagnostic or therapeutic use.