Protocol G0 Denaturation & Hybridisation Protocol

Combined Whole Chromosome Paints & Pan Centromeric Chromosome Paints

(Protocol G0)

XstarFISH[®]

Approx time:					
	Probe Preparation: 20 min + 16 hrs (Overnight) +30 min				
Solutions to be prepared:	20XSSC				

Solution 20XSSC:

87.6g NaCl <u>44.1g</u> Na Citrate up to 500ml Deionised Distilled water

Adjust pH to 7.0 using concentrated HCI (before finalising water volume), aliquot and autoclave. Store at 4° C).

Note: Ensure all solutions are mixed well.

Procedure: Slide pretreatment

- 1. Cells are fixed in 3:1 ethanol:acetic acid for 30 min, and then let them dry. In order to remove the acid which will interfere with the hybridization the slides are passed through a graded alcohol series 70, 90, 100% for 2 min. Air dry and then the slides can be stored for several days.
- 2. In order to ensure the cells stick to the slide the slides are baked at 65°C for 15 mins
- 3. After baking allow the slides to cool and then transfer slides to acetone for 10 mins
- 4. Air dry slides
- 5. In order to remove RNA which will cause non-specific binding of the probe and give background, incubate the slides for 1h at 37°C in 2X SSC+ RNase.(100µg/ml). Add 200µl per slide and cover with a parafilm coverslip. Incubate for 1hr at 37°C in incubator.
- 6. Wash off in 2X SSC for 5 min
- 7. Wash in PBS for 5 min
- Remove excess protein by incubating slides in Pepsin. In an Eppendorf tube pipette 1ml of 10mM HCl (stored at 4°C) and add 0.5µl of stock pepsin solution (1mg/ml). Vortex and add 200µl/slide in humidified slide chamber. Cover with parafilm coverslip and leave at room temperature for 2-3 mins.
- **Note: The timing is critical, do not overtreat**. Some slides will require different timings depending on the amount of cytoplasm. (NB some cells may require post fixing in formaldehyde after pepsin treatment to retain morphology).
 - 9. Wash slides in PBS for 5 min.
 - 10. Dehydrate through ethanol series.
 - 11. Air dry slides
 - 12. Denature cellular DNA by immersing slides in 70% formamide in 2X SSC at 70°C for 2 min. (15ml 2X SSC + 35ml formamide).
 - 13. After denaturation place slides in ice cold 70% ethanol for 2 min.
 - 14. Dehydrate cells by passing through ethanol series 2 minseach step.(70%, 90% and 100%).
 - 15. Air dry slides.

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Procedure: Probe preparation

- 1. Warm probes to 37°C for 5 min and mix well.
- 2. Add 3ml of whole chromosome paint to 12ml of hybridisation buffer in a microcentrifuge tube.
- 3. Denature probe for 10min at 65°C and hold at 37°C for 30-60 min.
- 4. Add 1ml concentrated centromeric probe + 5ml of hybridisation mix in a separate microcentrifuge tube.
- 5. Denature probe for 10 min at 85°C. Immediately chill on ice.
- 6. Combine whole chromosome probe and centromere probe and apply immediately to denatured cells on slide.
- 7. Hybridise for approximately 16 hours at 37°C in a humidified chamber.

Procedure: Post Hybridisation Wash

- 8. Remove cover slip washing for 5 min at 37°C in 2X SSC.
- 9. Wash slides twice in 50% formamide / 2X SSC 37°C, for 5 min each time.
- 10. Wash slides in 2X SSC, twice for 5 min each time.

Proceed directly to Detection step in Detection protocol for relevant labels. Do not duplicate Post Hybridisation wash.

Biotin Labelled Paints	
1043-KB-50	Detect using Cyanine 3
1596-KC-50	Detect using Cyanine 5
1082-KT-50	Detect using Texas Red
1089-KB-50	Detect using FITC

FITC Labelled Paints	
1084-KF-50	FITC Amplification Kit

Cyanine 3 Labelled Paints		
Cyanine 3-09	Direct labelling	

Dual Colour Painting	
1597-KD-50	Biotin-Cyanine 3 & FITC
1090-KD-50	Biotin-Texas Red & FITC
1084-KF-50	FITC & Cyanine 3

For detection kits and protocols please contact us or visit our web page at www.cambio.co.uk

This product is for research use only

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