

(1695-B-*) (1695-F-*)
(1697-B-*) (1697-F-*)

Approx time:

Probe Preparation: 10 min + 16 hrs (Overnight) +30 min

Solutions to be prepared:	20XSSC
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Solution 20XSSC: 87.6g NaCl
44.1g Na Citrate
up to 500ml Deionised Distilled water

Adjust pH to 7.0 using concentrated HCl (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 hybridisation 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
8. 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 overtrear.** 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 mins each step.(70%, 90% and 100%).
15. Air dry slides.

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Cambio Ltd, The Irwin Centre, Scotland Road, Dry Drayton, Cambridge, UK, CB23 8AR
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Procedure: Probe preparation

1. Warm probes to 37°C for 5 min and mix well.
2. Take 12.5µl of regular probe or 1µl concentrated probe per slide, plus hybridisation buffer to a total of 12.5µl per slide, and add to a microcentrifuge tube.

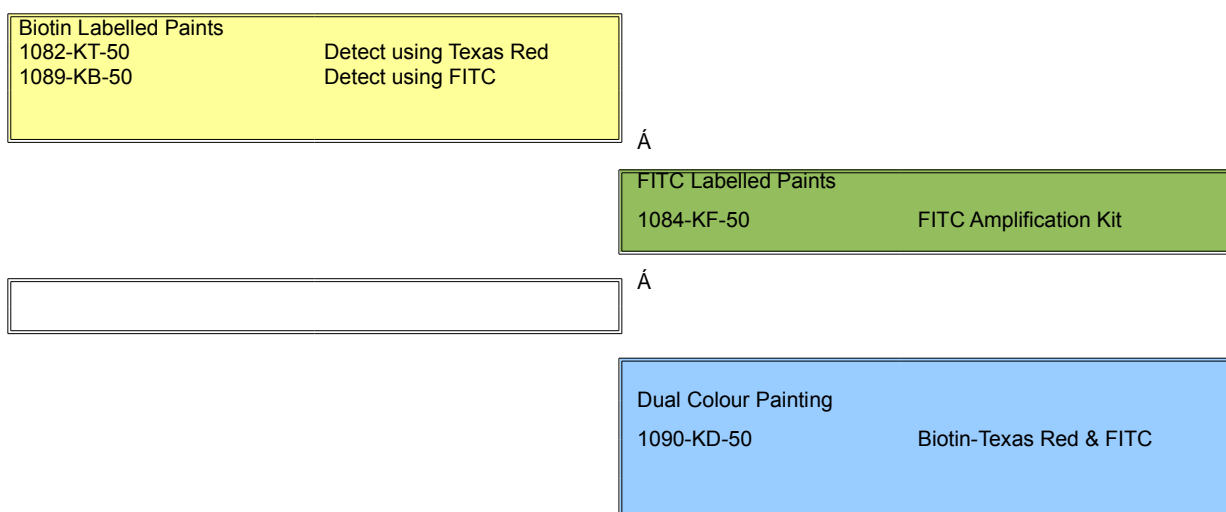
Note: *These amounts are based on a 22x22mm coverslip.
It is not recommended to use more than 3 concentrated probes in one 12.5µl mix.*

3. Denature chromosomes on slide in 70% formamide in 2X SSC for 2mins at 70°C. Immerse in ice cold 70% ethanol and dehydrate through a series of alcohol washes 70%,90% 100%. Dry at room temperature.
4. Denature probe for 10 min at 85°C. Immediately chill on ice.
5. Apply probe to slide.
6. Hybridise for approximately 16 hours at 37°C in a humidified chamber.

Procedure: Post Hybridisation Wash

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



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

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