# Biotin labelled chromosome Detection Protocol detect with Cy3



#### Introduction

The FISH protocol is divided into two stages. Denaturation and Hybridisation are performed on **Day One**. Washing and Detection are performed on **Day Two**.

On day one, the DNA of the chromosomes and paints are denatured and the hybridisation process (reannealing) takes place overnight. On day two, the slides are washed to remove unbound DNA sequences followed by detection, counterstaining and mounting.

Kit Contents				
Product Code	Description	Volume		
1124-Y1-50	Detect Y1 (Cyanine 3 Streptavidin)	20µl		
1124-DT-25	Detergent (Tween 20)	2 x 1ml		
1124-MD-50	Reagent MD (Antifade + DAPI)	2 x1.25ml		

#### **Requirements (not provided)**

Equipment	Reagents
Ethanol cleaned slides	Sodium Chloride
Coverslips	Sodium Citrate
Eppendorf tubes	HCI
Coplin jars	Formamide
Humidified chamber	Absolute Ethanol
Micro-pipette 1µl, 10µl, 500µl	Fixogum rubber cement
Pipette 10ml, 20ml	Clear nail varnish
Vortex	Deionised Distilled water
Parafilm	
Micro-centrifuge	
45°C Water bath	
37°C Incubator	
Fluorescence microscope with a suitable filter set	

#### Approx time:

Preparation 20 min Procedure 40 min

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Solutions to be prepared:

20XSSC 1XSSC 4XSSC Detergent wash solution Stringency wash solution Working Reagent A

Solution 20XSSC:	87.6g	NaCl		
	-	<u>44.1g</u>	Na Citrate	
		up to 5	00ml Deionised Distilled water	

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

Solution 1XSSC:	25ml	20XSSC <u>475ml</u> Deionised distilled water 500ml 1XSSC
Solution 4XSSC:	100ml	20XSSC <u>400ml</u> Deionised distilled water 500ml 4XSSC
Detergent wash solution:	500ml	4XSSC <u>250μl</u> Detergent DT 500ml Detergent wash solution
Stringency wash solution:	50ml	Formamide <u>50ml</u> 1XSSC 100ml Stringency wash solution
Stringency wash solution can be reused up to 5 times but should be discarded after 2 months		

Working Reagent A:2.5µlDetection reagent Y11247.5µlDetergent wash solution1250µlWorking Reagent A (Y1) (1:500)Incubate in the dark for 5 min. Microcentrifuge at 11.000g for 5 min.

Note: Ensure all solutions are mixed well.

All solution volumes sufficient for 10 slides

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### Procedure: Washing

- Pre-warm to 45°C in a water bath at least 30 min before starting: Two Coplin jars of Stringency wash solution (50ml each) Three Coplin jars of Solution 1XSSC (50ml each) One Coplin jar of Detergent wash solution (50ml)
- **Note:** The temperature is important. Check the temperature of the solutions in the Coplin jar and not of the water in the water bath.
  - 2. Take out the slide from the incubator and leave in Solution 1XSSC for 5 min. Take off rubber cement and replace in Solution 1XSSC to remove the coverslip.

**Note:** Do not allow to dry.

- 3. Stringency washes:
  - Wash slides twice by incubating 5 min each in Stringency wash solution (45°C). Wash slides twice by incubating 5 min each in Solution 1XSSC (45°C). Incubate slide for 4 min in Detergent wash solution. (45°C),

### **Procedure:** Detection

- 4. Apply 100µl of Working Reagent A onto the slide and cover with Parafilm immediately.
- 5. Incubate slide in a humidified box for 15-20 min at 37°C.
- 6. Remove Parafilm from the slide and wash 3 times for 4 min each time in the Detergent wash solution at room temperature, by emptying and refilling the Coplin jar.
- 7. Drain slide well and mount with 50µl of Reagent MD.
- 8. Apply glass coverslip and seal with nail varnish. Store slides in the dark at 4°C.
- **Note:** You get almost no air bubbles when Reagent MD is applied on the coverslip and the almost dry (but not dried out!) slide is laid face-down on the coverslip
  - 9. View slides using standard epifluorescence filters for Cyanine 3 and for counterstain DAPI.