Chromosome Preparation from Leucocyte Culture
A Simplified Method for Collecting Samples by Post

JANET M. ANDERS, ELIZABETH C. MOORES, and RICHARD EMANUEL

From the Institute of Cardiology and National Heart Hospital, London W.1

A convenient postal method for leucocyte culture was developed during the study of 136 cases of congenital heart disease (Anders, Moores, and Emanuel, 1965). The technique is a simplified version of that reported by Edwards (1963) and is based on his suggestion for the use of dextran to agglutinate the erythrocytes, which makes gravity separation quicker and leaves the majority of the leucocytes suspended in the plasma. Instead of defibrinating by shaking the blood with glass beads for 10 minutes, as described by Edwards (1963), we used heparin as an anticoagulant. Boots' preparation (Heparin B.P.) proved satisfactory for suppressing clotting throughout incubation. One of the disadvantages of heparin is that fibrin may precipitate during harvesting when hypotonic sodium citrate solution is added. In our experience, however, this does not occur if the original method is used, in which the culture medium containing the cells is diluted with three times its volume of water (Moorhead, Nowell, Mellman, Battips, and Hungerford, 1960). Finally, we found it unnecessary to remove the dextran by centrifuging through sucrose solution after hypotonic treatment (Edwards, 1963), if the cells were washed in several changes of fixative before making the slides.

Dextran was first used in this laboratory when difficulty had been experienced in separating plasma and leucocytes in polycythaemic patients with cyanotic congenital heart disease. Satisfactory separation was achieved in all cases when the whole blood was mixed with dextran, though in patients with a particularly high packed cell volume equal quantities of blood and dextran were necessary. It is now our standard practice for all cultures to add 2 ml. dextran to 10 ml. whole blood. When only small quantities of blood are available the same amount of dextran (2 ml.) can be added to as little as 1 ml. of blood with surprisingly good results.

Phytohaemagglutinin (Burroughs Wellcome) was used only as a mitotic stimulant and not for separating the plasma, as described in the standard method (Moorhead et al., 1960). It is therefore added immediately before incubation with the advantage that cultures may be kept at room temperature until a convenient day for incubation, making week-end harvesting unnecessary.

This technique has been adapted for the collection of samples by post. The method requires 10 ml. of blood from venepuncture: this is added to a bottle containing 0·1 ml. Heparin (Boots', 5,000 units per ml in normal saline) in 2 ml. dextran (Benger's Dextraven: 6% dextran in saline average molecular weight 150,000). After standing for 15-30 minutes the erythrocytes sediment and the supernatant plasma with leucocytes is then transferred with a fresh syringe and needle to a second bottle containing 5 ml. of tissue culture medium 199 (Glaxo). If the blood is left to stand for as long as one hour the leucocytes may also sediment, forming a fluffy coat, which makes separation difficult. It is possible to remedy this by shaking the blood and leaving it to settle again, though some of the leucocytes will be lost by this treatment. Both bottles are then posted to the laboratory where the culture is set up and incubated in the usual way.

A total of 26 samples has been obtained by post and 23 were cultured successfully. The average time between taking the blood and incubation was 3 days. The maximum delay was 5 days, when two cultures were mislaid on arrival at the hospital. One of these grew satisfactorily, the other failed. There was no apparent technical reason for the failure of the other two unsuccessful cultures.

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A postal pack for a single culture contains:

Bottle A: 0.1 ml. Heparin in 2 ml. dextran
Bottle B: 5 ml. T.C. medium '199'
2 disposable syringes (10 ml.)
1 disposable needle for taking blood
2 long needles for addition of blood to bottles
2 air vents
Alcohol and cotton wool for cleaning skin and bottles
Instruction sheet.

Bottles A and B can be kept in a refrigerator at 4°C. for two weeks. Storage for longer periods has not been investigated.

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References