Bedside control of heparin therapy by a simple whole blood clotting method

C. COTTON KENNEDY AND M. J. ROCKS  
From the Haematology Department, Belfast City Hospital, Belfast

Control of heparin therapy is most often monitored by whole blood clotting times. Based on the old-established method of Lee and White (1913) it is ideally carried out in the ward by a trained laboratory technician, the clotting process being allowed to occur in glass tubes, which are inverted at intervals and incubated in a portable water bath at 37°C.

In the past few years there has been an increase in the number of patients receiving heparin for a variety of conditions. Frequently laboratory control is not sought or is not available. When it is required it is often difficult in a busy laboratory to provide a technician to cover all the demands, especially at inconvenient times. In recent years we have developed a simple, accurate method which can quickly be demonstrated to the ward doctor who, without calling on laboratory help or using a water-bath, can thereafter measure clotting times at the bedside.

Method

In the ward two standard glass tubes (8 mm x 75 mm) are warmed in the hand for three minutes. Two ml venous blood is then withdrawn and a stopwatch is started as soon as the blood enters the syringe. One ml blood is dispensed into each of the two tubes. The tubes are held tightly in the hand and are inspected by tilting gently just beyond 90° every 30 seconds until the blood clots. The whole blood clotting time is taken as the average time of clot formation in the two tubes.

Hand temperatures measured in our series varied from 29°C to 33°C.

Results

To establish the normal range for the whole blood clotting time 100 normal subjects were selected.

The Lee and White method, using a water bath at 37°C, and the hand temperature technique were carried out simultaneously by two independent observers and the results were compared (fig 1).

Received for publication 8 August 1973.
heparin therapy, again using both whole blood clotting methods simultaneously. The results obtained are shown in figure 2.

**Comment**

Substitution of the hand temperature technique for that of the water bath method has enabled ward doctors, unused to performing coagulation tests, to measure accurately clotting times at the bedside.

We stress that brief instruction from trained laboratory staff should be given on the first occasion.

According to our findings hand temperatures, several degrees below 37°C, make little difference to whole blood clotting times. Based on our results of the Lee and White method at 37°C and the hand-held method at lower temperatures, we regard the normal whole blood clotting time by the latter technique to be between four and seven-and-a-half minutes.

We thank our medical and technical colleagues at the Belfast City Hospital for their cooperation and Miss E. Mullan, who typed this paper.

**Reference**


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**Letters to the Editor**

**Candida Infections**

The paper by Stieritz, Law, and Holder (_J. clin. Path.,_ 1973, 26, 405-408) states that all Candida isolates from patients should be speciated. Isolation of Candida species representing colonization of skin and mucous membranes occurs daily in most clinical laboratories. It is our experience that less than 1% of these are associated with infection with this genus. Although isolation from blood and urine is more suggestive of infection, many of these prove to be transient or spurious.

Continued study of the relative pathogenicity and antimicrobial susceptibility of Candida species is desirable. This may be accomplished in the majority of hospitals by referring isolates, which are repeatedly isolated from patients with symptoms suggesting infection, to reference laboratories which maintain the skill and materials required for accurate speciation. Despite acknowledged differences in the pathogenicity of Candida species, all may produce fatal systemic infections. The decision to treat should be based on clinical evidence of infection, not the species isolated. It is academically satisfying to have speciation but it is not essential to diagnosis and treatment. Laboratory workers and those responsible for licensure and performance evaluation may infer from these authors' remarks need and justification for new minimum laboratory standards. Increasing public reaction to rising health care costs dictates that we give greater attention to the cost benefit of arbitrary often academically inspired recommendations for increasingly expensive clinical microbiology.

RAYMOND C. BARTLETT
Division of Microbiology,
Department of Pathology,
Hartford Hospital,
Hartford, Conn., USA

**Klebsiella Species in Chest Infections**

I read with interest Dr Fallon's (_J. Clin. Path.,_ 26, 253) re-assessment of the significance of Klebsiella species in chest infections and note that he confirms the difference in this respect between _K. aerogenes_ and other species.

May I make the following observations?

1. Now that digestion-dilution methods are widely employed to avoid deceptive false positive cultures in sputum bacteriology (Wilson and Martin, 1972), I doubt the wisdom of using centrifuged deposits of sputum, other than for fungal isolations. This will have the effect of concentrating small numbers of coliforms, which will overgrow the mixed normal flora but would have been clearly identified as of doubtful significance by routine techniques. Though it may be quibbling, in many instances probably overemphasizes the importance of coliforms in sputum to talk of secondary invasion. Their invasive potentialities are nil in patients with normal immune mechanisms. Secondary colonization is probably a truer assessment of their significance.

2. I think that 10 years on one can no longer recommend a return to eponym titles such as Friedlander's bacillus. Perhaps a satisfactory compromise will be the term 'respiratory Klebsiella' used in the current edition of Topley and Wilson.

J. H. DARLING
Royal Postgraduate Medical School,
Hammersmith Hospital,
London

**Reference**