moglobin does not affect the level of alkali-resistant haemoglobin. The poor reproducibility experienced by Beaven et al (1960) when using cyanmethaemoglobin has not been encountered.

NORMAL VALUES
Estimations of 75 normal non-pregnant females and 24 normal males between 18 and 45 years are shown in Figure 2. An upper limit of 0·9% has been taken as normal because the distribution becomes discontinuous above this level.

References

Technical methods

A technique for demonstrating fibrinolysis by cutaneous bacteria

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Substances which convert plasminogen to plasmin, ie, activators of plasminogen, derive from various sources. They were first demonstrated in filtrates of haemolytic streptococci (Tillet and Garner, 1933) and since have been found in almost all fluids and organs of the body (Fearnley, 1965) and have been demonstrated in culture filtrates of some strains of staphylococci (Elek, 1959). Substances collected from the skin surface have also been shown to have fibrinolytic activity (Dawber, Nishioka, and Ryan, 1971) and it has been suggested that this activity, together with the clot-promoting effect of surface lipids, is important in reconstituting the skin surface following injury.

Dawber and his colleagues (1971) failed to find fibrinolytic activity among the bacteria of the normal flora of the skin. As some strains of Staphylococcus aureus have long been known to produce staphylokinase (Elek, 1959) it was decided to study skin isolates of these organisms and the closely linked groups, Staph. epidermidis and Micrococcus species. The investigation was later extended to include isolates of the cutaneous diphtheroids.

Materials and Methods

Fibrinolytic activity of bacteria has previously been demonstrated and measured using the fibrin plate technique (Astrup and Mullertz, 1952) in which a solution of fibrinogen is clotted with thrombin in a Petri dish and a small amount of the substance to be assayed placed on the fibrin so prepared. The plate is then incubated for a set period, usually about 20 hours, and lysis is shown as a clear zone which gives a measure of the fibrinolytic activity.

This technique is not ideal for testing large numbers of strains of bacteria and it was therefore modified and the tests performed in 50 × 6 mm test tubes. In each sterile test tube were placed 0·5 ml fibrinogen (1% in veronal buffer, pH 7·4), 0·02 ml of 0·1M calcium chloride, and 1 drop of peptone water. All solutions were sterilized by filtration before use. A large inoculum taken from an agar plate culture of the appropriate bacterial isolate to

Received for publication 1 March 1972.
be tested was added and finally 0·02 ml sterile bovine thrombin (0·5 units). Thorough mixing was ensured using a vortex stirrer (Griffin) and the resulting fibrin clot in which the test strain was suspended was incubated at 37°C and examined after four and a half hours, 20 hours, and then daily until control tubes showed evidence of clot disintegration. Bovine and human fibrinogen (Sigma) were used and the control fibrin clots were stable for more than four days in the case of the bovine fibrinogen and over three days for the human fibrinogen. The addition of the small amount of peptone water did not affect the stability of the clot but minimal medium produced fibrinolysis within 20 hours.

In all, 195 strains of _Staph. aureus_, 156 strains of coagulase-negative staphylococci and micrococci, and 110 isolates of cutaneous diphtheroids were examined.

Culture filtrates of a limited number of the strains (12) were also tested, and the results obtained with these were similar to those obtained with the organisms themselves.

**Results**

Fibrinolytic activity amongst the isolates tested is shown in Table I. There was no evidence of activity after four-and-a-half hours either using the organisms themselves or the culture filtrates. The phage types of the _Staph. aureus_ strains were known but the only significant difference between the various groups was the lower incidence of activity on bovine fibrinogen of the strains of phage group II (38% as compared to 87% of the strains belonging to the other groups). There was no significant difference in incidence of activity amongst the Baird-Parker _Staphylococcus_ and _Micrococcus_ subgroups ($\chi^2 = 8·71; 7$df) nor between the fluorescent and non-fluorescent diphtheroids tested ($\chi^2 < 0·01$). Two isolates of _Corynebacterium acnes_ were included but because of the difficulty of preserving a stable clot under anaerobic conditions, it was not possible to determine fibrinolytic activity.

**Comment**

Fibrinolytic activity is widespread amongst bacteria and can readily be demonstrated. The fibrin plate is a convenient method when only a few samples are being examined. However, there is a maximum of four or five specimens which can be assayed on one plate and even with this small number, difficulties arise when reading the results, for disintegration of the clot makes prolonged incubation impossible. Using test tubes, each strain may be separately tested for activity and the stability of the clot greatly improved; readings could easily be taken three to four days after inoculation. A quantitative assessment can be made by the time taken for lysis to appear.

**References**


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doi: 10.1136/jcp.25.8.740

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