Efficiency tests on a series of common skin antiseptics under ward conditions

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SYNOPSIS  Bacteriological trials were carried out on 11 different skin disinfecting agents to ascertain their relative efficiencies under ward conditions. A statistical analysis of the results is given. Of the antiseptics used, 5% Laurolinium appeared to be the most effective and has been adopted for routine use in the wards.

In 1958, a survey of sterilizing methods was made in two of the hospitals of the Portsmouth Group before the introduction of a central sterile supply.

It was found that 60% of the sterile procedures carried out in the wards were the giving of hypodermic injections and that the techniques and skin disinfectants used varied considerably from one ward to another. This paper describes the investigations carried out to determine the efficiency of skin disinfection before injection under ward conditions.

HISTORICAL

In the past many of the tests made to evaluate skin disinfection were of a laboratory nature. Fleming in 1924, using the Wright 'slidecell' method, illustrated the germicidal and leucocyte-inactivating properties of blood and used it for evaluating a number of antiseptics. Another approach was to study the effect of antiseptics on the skin of animals in vivo and on egg membranes (Browning, 1934; Hunt, 1937; Green and Birkeland, 1944). These tests, however, were of little practical value in assessing the effects of disinfectants on human skin.

In his book 'Disinfection and Sterilization,' Sykes (1958) described a direct swabbing technique which he had found satisfactory. This tested the retention of activity by disinfectants other than soaps and had been adapted from that described by Colebrook and Maxted (1933). A similar method was proposed by Story (1952) in which small circles of skin on the forearm were infected with a drop from a chosen culture. Various handwashing tests have been published by different workers devised primarily for testing antiseptic soaps and, of these, that described by Cade (1950) seems to be most generally used. None of these investigations gave any indication as to the number of bacterial colonies likely to be found on the skin of patients, nor of the ability of disinfectants under ward conditions to reduce their numbers. In addition they did not simulate the actual procedure carried out by a nurse when swabbing the arm of a patient before giving an injection. In view of the fact that the nurse swabs the area of skin once or twice at the most, it was felt important to determine whether any of the antiseptics used in this way were effective.

Eleven antiseptics were chosen for trial, since they appeared to be those most commonly used in the hospitals where the tests were carried out. These were: Roccal 1 in 40, methylated ether, cetrimide 0.5% with Hibitane, 0.05% Penotran, 1 in 20 dequalinum chloride, 0.5% merthiolate in aqueous solution, 1 in 1,000 tinct. merthiolate, 1 in 1,000 with 50% alcohol, solution of iodine 1% in 90% spirit, and Laurolinium 1% and 5%.

METHOD

To determine the number of organisms on a given area of skin the technique of 'replica plating' was used (Lederberg and Lederberg, 1952). Replica plating was also used to provide a simple form of viable count by Elek and Hilson (1954). The advantages of this method are that a relatively large area of skin, 1 in. x 1 in., can be tested and the transfer carries over very little more antiseptic than the amount adhering to individual organisms. The replica blocks used consisted of 1 in. x 1 in. x 1 in. wooden blocks on one of whose surfaces was glued a square of velvet 1 in. x 1 in. This provided the 'contact' area. One of these blocks could be used many times and was easily sterilized by steam. The portion of skin used for the tests was the same for every patient, namely, an area on the inner aspect of the arm 2 in. below the cubital fossa. The culture media used were nutrient agar for methylated ether, 1% iodine in 90% spirit, and 70% alcohol, and blood agar for all the other disinfectants.
To carry out the trial, the block was first placed on the area B (shown below) of a culture medium plate to pick up a thin film of agar on its velvet surface. The block was then applied to the patient's arm to pick up bacteria and then pressed lightly back onto the same area of the plate. The arm was then swabbed quickly twice with a ball of sterile cotton wool dipped in the antiseptic solution being tested. Another identical sterile 'replica' block was taken and placed on square A of the agar plate. It was then pressed lightly first onto the patient's arm over the swabbed area and then onto the same square.

![Diagram of an agar plate used in the test. One plate was used for each two patients.](image)

This procedure was repeated on 100 different patients, male or female, for each solution under trial. After 24 hours' incubation at 37°C, the plates were examined and the number of colonies of bacteria on each square counted.

Certain possibilities have to be considered when carrying out these tests. Since the area of skin used for each of the two samples was the same, difficulty might have been experienced when interpreting the results because the bacterial population might, in fact, have been reduced by the first sampling, and the apparent reduction in colony count be due to this factor and not to the action of the particular antiseptic solution used in the test. In order to exclude this possibility a further 100 tests were carried out in which the same skin area was sampled twice without any intervening swabbing with antiseptic solution.

A further possibility also had to be taken into account when interpreting the results and that was 'carry-over' of the antiseptic fluid to the culture plate. Should this happen the counts would not reflect truly the immediate disinfecting powers of the solutions. This applies particularly to the slower-drying aqueous solutions.

Supplementary investigations were, therefore, carried out on those solutions which were slower to volatilize by using culture medium to which a suitable inhibitor had been added, thus preventing continued action by any of the antiseptic carried over. These supplementary tests were carried out on both of the Laurolinium solutions and the cetrimide-Hibitane mixture, since from early results it appeared that these solutions had not completely evaporated before the skin was resampled after swabbing. The inhibitor used was 1% lubrol W with 0.5% lecithin.

For each of these tests four blocks and two blood agar plates, to one of which the inhibitor had been added, were used. Samples were taken simultaneously from two adjacent sites on the routine test area of the arm and platted out onto square B of the two different culture plates. Both skin areas were then swabbed with the test solution and a further two blocks applied to them and again pressed onto the two areas A of each plate.

**RESULTS**

It was found that taking two consecutive samples from the same area of untreated skin without intervening swabbing slightly reduced the number of colonies isolated from the second replica block when compared with the first, but the reduction was not statistically significant. The counts on inhibitor and non-inhibitor culture plates also showed no significant difference. These two possibilities can, therefore, be discounted when assessing the relative efficiency of the different disinfecting fluids.

The table summarizes the results of the plate counts before and after the application of the various fluids to the skin and gives their statistical significance.

Because the number of colonies counted showed such a wide range from 0 to more than 100, with the smaller counts predominating, the actual figures obtained were replaced by their square roots, as shown in column 3 of the table. These figures were used in the analysis. For example, if the number of colonies counted was 100 or more, the square root was taken as 10. This gave a more even distribution and made the statistical analysis more likely to be correct. The figures in column 3 show the mean decrease in terms of the square root of the number of colonies after swabbing. The antiseptic solutions are placed in the order of their apparent effectiveness.

**DISCUSSION**

The disinfecting agents fall into five groups, showing different degrees of effectiveness. Within the individual groups the solutions appear fairly uniform in their action.

The tests of significance given in columns 4 and 5 of the Table show that with the exception of Penotran and aqueous merthiolate (group I) all the solutions had a significantly destructive effect on the skin flora. Methylated ether (group II), though better than the group I reagents, was relatively ineffective. Dequalinium chloride and 70% alcohol (group III) were significantly more potent than the group II solutions.
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TABLE
RESULTS OF COMPARISON OF 11 SKIN DISINFECTING AGENTS

<table>
<thead>
<tr>
<th>(1) Skin Sterilizing Agents</th>
<th>(2) No. of Patients</th>
<th>(3) Mean Decrease in Square Root of No. of Colonies after Swabbing</th>
<th>(4) Statistical Significance of Decrease Compared with that for No. Agent</th>
<th>(5) Statistical Significance of Decrease Compared with that for Last Preceding Agent</th>
</tr>
</thead>
<tbody>
<tr>
<td>GROUP I</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>None (control test)</td>
<td>100</td>
<td>0.18</td>
<td>Not significant (P &gt; 0.10)</td>
<td></td>
</tr>
<tr>
<td>Penotran stock solution 1:20</td>
<td>100</td>
<td>0.29</td>
<td>Not significant (P &gt; 0.10)</td>
<td></td>
</tr>
<tr>
<td>Merthiolate (aqueous) 1 in 1,000</td>
<td>100</td>
<td>0.34</td>
<td>Not significant (P &gt; 0.10)</td>
<td></td>
</tr>
<tr>
<td>GROUP II</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methylated ether</td>
<td>100</td>
<td>0.82</td>
<td>Very highly significant (P &lt; 0.001)</td>
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<tr>
<td>GROUP III</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Dequalinium chloride 0.5%</td>
<td>100</td>
<td>1.47</td>
<td>Very highly significant (P &lt; 0.001)</td>
<td>Significant (P &lt; 0.05)</td>
</tr>
<tr>
<td>Alcohol 70%</td>
<td>100</td>
<td>1.56</td>
<td>Very highly significant (P &lt; 0.001)</td>
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<tr>
<td>GROUP IV</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Iodine 1% in spirit 90%</td>
<td>100</td>
<td>2.02</td>
<td>Very highly significant (P &lt; 0.001)</td>
<td>Not significant (P &gt; 0.10)</td>
</tr>
<tr>
<td>Merthiolate (tincture) 1 in 1,000</td>
<td>100</td>
<td>2.18</td>
<td>Very highly significant (P &lt; 0.001)</td>
<td></td>
</tr>
<tr>
<td>Cetrimide 0.5% and Hibitane 0.05%</td>
<td>100</td>
<td>2.1</td>
<td>Very highly significant (P &lt; 0.001)</td>
<td></td>
</tr>
<tr>
<td>Roccal (benzalkonium) 1 in 40</td>
<td>100</td>
<td>2.22</td>
<td>Very highly significant (P &lt; 0.001)</td>
<td></td>
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<tr>
<td>GROUP V</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Laurolinium 1%</td>
<td>100</td>
<td>3.16</td>
<td>Very highly significant (P &lt; 0.001)</td>
<td>Highly significant (P &lt; 0.01)</td>
</tr>
<tr>
<td>Laurolinium 5%</td>
<td>100</td>
<td>3.38</td>
<td>Very highly significant (P &lt; 0.001)</td>
<td></td>
</tr>
</tbody>
</table>

P = statistical significance of decrease.

The agents in group IV were not significantly more active than the 70% alcohol of group III, but the two Laurolinium solutions in group V were much the most efficient of all the skin disinfecting agents tested.

It is appreciated that we have no definite proof that lubrols W and lecinith will inhibit Laurolinium in the concentrations used for the trials.

In view of these findings, 5% Laurolinium has now been adopted as the standard pre-injection skin disinfectant in all wards in the Portsmouth Group which are served by the Central Sterile Supply Department.

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REFERENCES