In-use testing of disinfectants in hospitals

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SYNOPSIS One hundred and five samples of clear, soluble phenolic disinfectants were obtained from varying sites in the wards of six hospitals. The concentration of disinfectant in each sample was measured by a colorimetric method and bacterial contamination was measured by an ‘in-use’ test and a membrane filter technique. The concentrations of disinfectant in 24/105 (23%) samples were at the recommended level and 53 (50-5%) were below. Bacterial contamination with Gram-negative bacilli was found in 26/49 (53%) samples containing less than 0-8% of disinfectant and 5/86 (5-9%) samples containing more than 0-8%. The concentrations in two of the heavily contaminated samples were 1-5% and 1-6% respectively.

The wide variety of disinfectants used in hospitals has been described in several surveys (eg, Public Health Laboratory Service Report, 1965; Ayliffe, Brightwell, Collins, and Lowbury, 1969). Since these reports were published, disinfectant policies have been introduced in many hospitals, but others either have no policy or still use expensive and often inappropriate disinfectants for treating the environment. The principles of formulating a policy were described by Kelsey (1970), and the ‘in-use’ dilutions are usually chosen from the manufacturers’ recommendations or on the basis of the capacity test (Kelsey and Sykes, 1969). However, a laboratory test cannot reproduce the wide range of conditions which exist when the disinfectant is in use, and it is, therefore, advisable to carry out in-use tests for bacterial contamination (Kelsey and Maurer, 1966) when a new disinfectant is introduced into a hospital and at intervals afterwards. The in-use test will not determine whether contamination is due to an inadequate concentration of disinfectant or whether organisms are surviving or growing at or above the recommended concentration; inadequate concentrations of disinfectant in the absence of bacterial contamination will also not be detected. In this study, concentrations of disinfectants were measured under in-use conditions by means of a colorimetric test and bacterial contamination was assessed by an in-use test (Kelsey and Maurer, 1966) and by a technique using a membrane filter.

Methods

Samples were collected from six hospitals, all of which have a disinfectant policy and mainly use a clear, soluble phenolic disinfectant, Stericol, for environmental disinfection. One hundred and five samples of solutions of the phenolic disinfectant were collected from mop buckets, toilet-brush holders, thermometer holders, containers for contaminated instruments, and other in-use situations. Some samples of chlorhexidine, Savlon, and other disinfectants were also collected.

COLORIMETRIC MEASUREMENT OF CONCENTRATION OF PHENOLIC DISINFECTANT

Of 2% aminophenazone, 0-1 ml and 9-8 ml of 0-025% sodium carbonate were added to 0-1 ml of the disinfectant. Then 0-1 ml of 2% potassium ferri-cyanide was added, and after mixing readings were made in a spectrophotometer at 545 nm. A standard curve was prepared from concentrations of the same disinfectant treated in the same way as the sample. The purple colour obtained is reasonably stable but should be read within one hour. If no spectrophotometer or colorimeter is available, an approximate assessment of the concentration can be obtained by comparing the colour of the test and standard solutions with the naked eye.

BACTERIOLOGICAL CONTAMINATION

Direct culture method

Five drops (0-02 ml per drop) from a standard pipette of the sample were placed on a well dried plate containing nutrient agar and 5% horse blood and incubated for 24 hours at 37°C. A 1/10 dilution of the disinfectant in lecithin-Tween 80 broth was made before the above procedure was carried out.
when chlorhexidine or quaternary ammonium compounds were examined. Counts were made of the number of colonies grown from the five drops. Further incubation of the plates or dilution of the disinfectant did not appear to influence the results with phenolic disinfectants in this study, but for routine testing, incubation for 72 hours at 37°C and at room temperature is advisable.

**Membrane filter method**

One ml of the disinfectant solution was filtered through an Oxoid membrane filter (pore size 0.45μm) using a millipore sterilfil filtration system. The filter was washed through with 20 ml of broth, containing a neutralizer if necessary, and then transferred to a 5% blood agar plate. After incubation at 37°C for 24 hours, counts were made of the colonies growing on the membrane. Gram-negative bacilli were identified by the method of Cowan and Steel (1965).

**Results**

Table I shows that only 24/105 of the samples were at the recommended concentration and many were much higher when measured by the colorimetric method. Some form of measurement was used in all hospitals except hospital 6. The concentrations found in hospital 5 were more accurately measured than the others, since disinfectant was issued to the ward diluted ready for use.

**Bacterial contamination of samples**

A sample was considered to be contaminated if more than 50 organisms/ml were grown, either on the membrane filter or in the in-use test, and using this criterion the results from both methods were similar; most contaminated samples contained more than 1 000 organisms/ml. Table II shows that most samples containing less than 0.1% of disinfectant were contaminated (11/13, 84.6%); in samples containing between 0.1 and 0.8% of disinfectant, 15/36 (41.7%) were contaminated, and when the concentration was between 0.8 and 2.0%, 5/34 (14.7%) were contaminated. None of the samples containing over 2.0% disinfectant was contaminated. The contaminating organisms were mainly *P. aeruginosa* or non-fermenting Gram-negative bacilli. At very low concentrations of disinfectant, Klebsiella sp. and *Escherichia coli* were commonly found, especially in mop buckets; spore-bearing bacilli were not included in the results. Table III shows the site of contamination and concentration of disinfectant of the five samples with a concentration at or above that recommended. All were contaminated with non-fermenting, non-pigmented, oxidase-positive, Gram-negative bacilli. On sub-

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### Table I: Variation in concentration of clear soluble phenolic disinfectant

1Hospitals 1-4, 6, recommended concentration 1%, range 0.8-1.2%; Hospital 5, recommended concentration 2%, range 1.8-2.2%.

<table>
<thead>
<tr>
<th>Hospital</th>
<th>Total No. of Samples</th>
<th>No. Below Recommended Concentration</th>
<th>No. at Recommended Concentration</th>
<th>No. Above Recommended Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>15</td>
<td>8</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>2</td>
<td>9</td>
<td>5</td>
<td>0</td>
<td>4</td>
</tr>
<tr>
<td>3</td>
<td>32</td>
<td>17</td>
<td>7</td>
<td>8</td>
</tr>
<tr>
<td>4</td>
<td>11</td>
<td>10</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>5</td>
<td>17</td>
<td>5</td>
<td>10</td>
<td>2</td>
</tr>
<tr>
<td>6</td>
<td>21</td>
<td>8</td>
<td>3</td>
<td>10</td>
</tr>
<tr>
<td>Totals</td>
<td>105</td>
<td>53</td>
<td>24</td>
<td>28</td>
</tr>
</tbody>
</table>

### Table II: Contamination and concentration of clear soluble phenolic disinfectant

<table>
<thead>
<tr>
<th>Equipment in Disinfectant Solution</th>
<th>Percentage Concentration of Disinfectant</th>
<th>No. of Organisms/ml in Disinfectant Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sink mop</td>
<td>0.95</td>
<td>Uncountable</td>
</tr>
<tr>
<td>Floor mop</td>
<td>0.9</td>
<td>50</td>
</tr>
<tr>
<td>Floor mop</td>
<td>1.5</td>
<td>Uncountable</td>
</tr>
<tr>
<td>Floor mop</td>
<td>0.9</td>
<td>Uncountable</td>
</tr>
<tr>
<td>Cloth for surface cleaning</td>
<td>1.6</td>
<td>Uncountable</td>
</tr>
</tbody>
</table>

### Table III: Contamination of phenolic disinfectant solutions at concentrations above 0.8%
culture in nutrient broth these strains were inhibited by a concentration of 0·5% of disinfectant.

**Discussion**

Contamination of disinfectants was satisfactorily assessed by a simple in-use test (eg, Kelsey and Maurer, 1966); the membrane filter technique, although more accurately detecting small numbers of organisms, showed little advantage in testing phenolic disinfectants in the present study. It is probable that the membrane filter technique would be more useful for sampling solutions of chlorhexidine or quaternary ammonium compounds, but contamination of these compounds was too low in this study for assessment.

A simple colorimetric test for measuring the concentration of a phenolic disinfectant is a useful addition to the existing bacteriological tests. However, it should be emphasized that the chemical estimation described cannot replace the bacteriological in-use test for control purposes, since the activity of a disinfectant depends on formulation as well as phenol content. The study showed that measurement of disinfectants was inaccurate and that contamination was usually due to inadequate concentration, ie, of 105 samples examined 26/49 (53%) were contaminated when the concentration was less than 0·8% and only 5/56 (8·9%) when the concentration was at or above this concentration. The isolation of organisms in large numbers in two of these samples containing 1·5% and 1·6% disinfectant respectively was clearly not due to inactivation of disinfectant, or to a recent addition of organisms since samples were not usually cultured until one to two hours after collection. The organisms were able to survive at a concentration which was usually rapidly bactericidal, but the property was rapidly lost, since on subculture the organisms were killed by 0·4 to 0·5% of disinfectant and resembled normally sensitive organisms. These strains may have become adapted to the higher phenolic concentrations or were possibly protected by a layer of protein which was lost on subculture. Adaptation of *Pseudomonas cepacia* to a 1 in 30 dilution of Savlon has been described (Bassett, 1971) and tolerance of the two organisms described here was increased again in laboratory experiments by habitation in increasing concentrations of the phenolic disinfectant to 1·0%, but not as yet to 1·5%. Contamination of diluted disinfectant in stock bottles was not found in the small number of samples examined in this survey, but there is always a risk of organisms surviving and possibly growing if containers are not washed and preferably disinfected before refilling.

The concentration of the disinfectant (1%) recommended for treatment of light contamination on the basis of the Kelsey-Sykes test is obviously adequate for most purposes, including floor mops, which might be considered heavily contaminated. Most of the hospitals visited found it easier to use one concentration (1%) although as indicated the concentration found in practice was very variable. The use of 2%, the concentration recommended for heavy contamination, might prevent all contamination, but would double the expense, increase the possibility of skin reactions, and it is still possible that organisms would become tolerant to the higher concentration. The use of a 1% solution and regular in-use testing (eg, two to four times a year) should detect the appearance of resistant organisms and perhaps a policy involving rotation of types of disinfectant should be considered; the possibility of cross-resistance between disinfectants requires further investigation. A more satisfactory answer would be to reduce the use of disinfectant solutions in hospital. Although equipment immersed in disinfectant solution was found less often in hospital wards during this survey than in earlier surveys, inappropriate uses of disinfectants were still commonly found. Many of the samples with inadequate concentrations were obtained from floor mop water or water for washing surfaces, where the use of a disinfectant was usually unnecessary. However, mops do require disinfection after use and heat treatment is preferable to chemical disinfection (Colquitt and Maurer, 1969).

If a disinfectant is used at all the concentration should be adequate, and a reliable method of measurement of disinfectant and diluent should be available. In hospital 5, disinfectant was issued to the wards at the correct use-dilution. Although clearly marked ‘use undiluted’, ward staff frequently added a ‘cupful’ to a bucket; five samples showing inadequate concentrations were found in this hospital. A further disadvantage of this method is that a considerable volume of water is unnecessarily transported from the pharmacy to the wards. In hospital 4, bottles containing a measured amount of undiluted disinfectant were issued to the wards. Although this method should have been satisfactory 10/11 samples examined showed concentrations below that recommended. Domestic staff often used half a bottle or less, rather than the whole bottle for reasons of economy. The amount of water in the bucket was rarely measured although the required level of water should have been marked on the interior of the bucket. Hospitals 1, 2, and 3 used a dispensing pump attached to a container of undiluted disinfectant. Concentrations were very variable and obviously this technique was not as
satisfactory as might be expected. Whichever technique of measurement is used, education of the domestic staff is obviously of major importance. A disinfectant policy will only work if it is understood and adhered to by all grades of staff. It would also be of value if policies involving other disinfectants were similarly assessed by chemical and bacteriological in-use tests.

We wish to thank the staff of the six hospitals for their cooperation.

References

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