Bacteriostatic and bactericidal actions of boric acid against bacteria and fungi commonly found in urine

P D Meers, C K Chow

Abstract

Boric acid has been used for over 20 years to preserve urine while in transit for bacteriological examination. It has been suggested that it may be toxic for some urinary pathogens. To investigate this several strains of bacteria and fungi commonly found in urine were exposed to different concentrations of boric acid in nutrient broth. Viable counts were made at the outset and at intervals for up to 24 hours at room temperature to detect bacteriostatic or bactericidal effects. At concentrations between 10 and 20 g/l boric acid was bacteriostatic or fungistatic for very nearly all the common urinary pathogens. At 10 g/l boric acid was weakly bactericidal for some strains of Acinetobacter calcoaceticus and Pseudomonas aeruginosa, though higher concentrations were bacteriostatic only. Group B streptococci varied in their response to boric acid, but for most of them 10 or 20 g/l was satisfactorily bacteriostatic. It is concluded that boric acid is rarely toxic, and when it is, the effect is usually sufficiently delayed to be of only theoretical importance.

In 1969 Porter and Brodie described the use of boric acid at a concentration of 18 g/l to preserve urine while in transit for bacteriological examination. Although urine is a culture medium, they showed that when boric acid was added the number of any bacteria present did not change significantly for up to 48 hours at room temperature, and other cellular elements remained substantially intact. They claimed this effective bacteriostasis allowed the numerical criteria for the laboratory diagnosis of urinary tract infection (UTI) to be applied despite delay in performing the examination. This cheap and simple way to increase accuracy in the diagnosis of bacteriuria and pyuria has not been widely adopted. Ten years after the introduction of boric acid for the preservation of urine only a quarter of a group of 64 laboratories in England and Wales were using it.

The relative lack of interest is perhaps explained by the small number of publications that describe the use of boric acid, and the suggestion in some of them that it is occasionally bactericidal. As toxicity and concentration are usually related and as we could find no adequate documentation for the choice of the 18 or 20 g/l concentration normally used, we exposed several strains of bacteria and fungi commonly found in urine to different concentrations of the acid in nutrient broth.

Methods

The culture media used were brain heart infusion (BHI) broth, cystine, lactose, electrolyte deficient (CLED) agar, and 5% horse blood agar. Dilutions of bacteria were made in quarter strength Ringer's solution (CM225, CM301, CM375, BR52 from Oxoid Ltd, Basingstoke, England). The boric acid powder came from BDH Chemicals Ltd, Poole, England. The bacteria and fungi tested had been isolated from clinical specimens (table). They had been identified by standard methods. The hospital laboratory in which the organisms were isolated does not receive specimens from sources in the community. Those available for testing had been taken from hospital patients.

Three linked experiments were performed. In the first, 89 sets of six glass bottles containing 10 ml volumes of BHI with boric acid at concentrations of 0, 2.5, 5, 10, 20 and 40 g/l were used. At 40 g/l boric acid is approaching the limit of solubility in water at room temperature. Each set of bottles was inoculated

<table>
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<tr>
<th>Bacteria and fungi</th>
<th>From urine</th>
<th>From other sites</th>
<th>Totals</th>
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<tr>
<td>Totals</td>
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<td>102</td>
<td>191</td>
</tr>
</tbody>
</table>

Varieties, numbers, and origins of bacteria and fungi tested
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with a culture of one of the bacteria that had originated in urine (table). The inoculum was prepared from a 24 hour culture in BH1 broth of the strain concerned, diluted first to match an opacity standard approaching a viable count of 10^5/ml. Further dilutions were made in quarter strength Ringer’s solution until an inoculum of 100 μl added to each bottle contained about 10^5 bacteria, to yield a target initial count in the broth of 10^4/ml. Immediately after inoculation 10 μl of broth from the BH1 bottle with no boric acid was spread on to agar, using blood agar for group B streptococci and CLED for the remainder. The plate was incubated at 37°C overnight and a colony count made to measure the inoculum. After the sets of bottles had stood on the bench at about 24°C for 24 hours viable counts were performed on all of them by the method described. The target viable count and the small transfer volume of 10 μl were chosen both to be of relevance in the diagnosis of UTI and to yield adequate sensitivity with easily counted colonies.

The second experiment involved bacteria and fungi originating in specimens other than urine. This was done to show whether the microbes from urine used in the first experiment had been selected by prior exposure to the boric acid, adopted as a preservative following earlier work. The second experiment was done in the same way as the first, though only the 0, 10, and 20 g/l concentrations of boric acid were inoculated with each organism. The inoculum for fungi was established by trial and error. Each set of three bottles was duplicated using inocula prepared from BH1 broth cultures that had been incubated for four and 24 hours. This was to see if bacteria or fungi in the logarithmic and stationary phases of growth differ in sensitivity to boric acid.

The third experiment involved single strains of Acinetobacter, Alcaloeceticus, Pseudomonas aeruginosa, and Streptococcus group B for which boric acid had proved somewhat toxic at one or more concentrations. Twenty four hour broth cultures of these were used as inocula, as for the first experiment, but subsequent viable counts were made hourly for the first six hours as well as at 24 hours.

For direct comparison the microbial counts observed were standardised to a common base. They were multiplied by a factor derived by dividing the target initial microbial count of 10^4/ml by the inoculum actually achieved in each set of bottles, measured in the control broth immediately after inoculation. Confidence limits for the precision of the counts were estimated by plating 20 replicate 10 μl volumes from each of four suspensions containing about 10^4/ml viable Escherichia coli, Staphylococcus aureus, group B streptococci and Candida albicans, respectively. The factors used to standardise these counts for inclusion in the figure were derived by dividing the 10^4 target by the means of each set of 20 counts.

Results

Preliminary experiments showed that growth was inhibited when 100 μl volumes of broth containing higher concentrations of boric acid were plated on to solid media. They also showed that heavily capsulated strains of Klebsiella spp gave inconsistent viable counts whether or not boric acid was present. The large mucoid colonies these bacteria produce on agar are assumed to arise from microcolonies of varying size that develop in liquid growth media where they are held together by capsular material. Each microcolony, spread out when transferred to a solid surface, produces a single large colony. In such circumstances a colony count underestimates the viable count by a variable and potentially large factor, so only non-mucoid strains were used in the final experiment.

Viable counts standardised to a common base are shown in the figure, which depicts the responses of all the microbes listed in the table to varying concentrations of boric acid. The graphs combine data from the first experiment with those from the second in which the inocula had also been pre-incubated for 24 hours. This accounts for the greater number of points at the 10 g/l and 20 g/l concentrations. When a comparison was made no difference was noted between strains that had and that had not previously been exposed to boric acid. The bars denoting confidence limits were constructed to take account of the fact that each point in the graphs resulted from two independent viable counts. The range observed experimentally was extended by multiplying the standardised values for the highest and lowest value in each case by itself, and dividing the product by the target count of 10^4. Control broths with no boric acid gave confluent growth when they were subcultured after 24 hours at room temperature, indicating counts in excess of 10^6/ml.

The second experiment yielded two pairs of viable counts for each microbial strain exposed to boric acid initially either in its logarithmic (four hour) or stationary (24 hour) phase of growth, at concentrations of 10 g/l and 20 g/l. The four and 24 hour counts were compared at each concentration by the Wilcoxon signed rank test. As the different enterobacteria responded so similarly they were treated as a group for this purpose. The other organisms were tested within individual genera or species. The closest approach to a significant difference was found with group B streptococci exposed to boric acid at 20 g/l (t = 2, p = 0.1). In the third experiment bacterial strains selected as being sensitive to boric acid were exposed to it at two concentrations and sequential viable counts were made. A viable count of Ps aeruginosa were observed to fall below the lower confidence limit (to 30% of the initial count) was after 24 hours exposure at 10 g/l. When A calcoaceticus was exposed to 10 g/l the count began to fall significantly at the fifth hour, and by 24 hours it had been reduced to 13%, of the initial level. At 20 g/l the count had not changed significantly at six hours, but it had fallen to 37% at 24 hours. Counts of group B streptococci had fallen to 36% and 42% of the numbers originally present in 10 g/l and 20 g/l boric acid by the second hour. Subsequent exposure for up to 24 hours produced no further reduction.
Dose-response curves of the bacteria and fungi listed in the table against boric acid, at room temperature. Counts are standardised to an inoculum of 10^4/ml (dotted lines) at the beginning of exposure. Confidence limits (bars) are derived from replicate counts of representative organisms.

Graph a, Escherichia coli; b, Klebsiella spp; c, Enterobacter and Citrobacter spp; d, Proteus, Morganella, and Salmonella spp; e, Pseudomonas aeruginosa — o, Candida albicans — o; f, Staphylococcus spp; g, Acinetobacter calcoaceticus — o, A lucofis — o; h, Streptococcus group B — o, group D — o.

Discussion

Bacterial multiplication in urine while in transit to the laboratory leads to cumulatively false positive results over time. Preserving with boric acid, refrigeration, dip-inoculation methods and limitation of delay before examination have been suggested as ways of overcoming this. Refrigeration can be inconvenient, and its effect is delayed by the time needed for a specimen to cool. It is also not easily applied over a distance. Dip-inoculation methods are popular, but the small surface area provided for interpretation can cause difficulty, particularly with mixed cultures. Their use generally precludes microscopy for the diagnosis of pyuria. Progressive desiccation of agar gels packed in plastic containers leads to an artificial increase in bacterial counts estimated by dip-inoculation, and thus increasing numbers of false positive results. Preservation with boric acid has none of these disadvantages.

So far as delay before examination is concerned, it has been suggested that no special precautions are necessary if urine is cultured within four hours of its production. This advice arose from an experiment in which specimens were collected from selected subjects, some of them hospital staff. Patients with glycosuria or urinary tract abnormalities were excluded, and there is no evidence that any of the specimens came from infants, children, or the aged. It might be deduced that the four-hour safety margin was assessed on specimens occupying the least heavily contaminated end of the normal spectrum. These would be expected to last longer before they yielded false positive results. In an experiment in which unselected routine clinical specimens were tested the “safe” period was found to be 90 minutes. This was less than the mean time taken for specimens to reach the laboratory in the hospital concerned, which had a collection service of not less than average efficiency. By the time they were examined unpreserved samples gave 5%, more cultures interpreted as positive than the preserved ones. In these circumstances some method for preventing the multiplication of contaminating bacteria or fungi should be applied to all urine specimens that are sent for bacteriological examination.

We have not found an account of the mechanism by which boric acid acts to produce bacteriostatic or bactericidal effects. Unlike some other antimicrobial substances it is not
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more toxic for rapidly growing than for stationary phase cells. For many microbes exposed for 24 hours, boric acid seems to have a dose-response curve that is long and nearly flat. This was particularly so for Ps aeruginosa, which displayed almost complete bacteriostasis over the range of boric acid concentrations used for all but one of the 17 strains tested. For the enterobacteria the flat range extended from 10 g/l to 40 g/l, with significant but incomplete bacteriostasis evident at 2.5 g/l and 5 g/l. A liquefying Ps aeruginosa, but most strains of A calcoaceticus were insensitive to boric acid up to 5 g/l. A partial bactericidal effect for some strains was noted at 10 g/l, but most higher concentrations were bacteriostatic. It may be that counts of these capsulated bacteria were unreliable for the same reason as suggested for Klebsiella spp.

All the members of the genus Staphylococcus reacted to boric acid much like Ps aeruginosa, but the dose-response curve for streptococci was steeper. Under the conditions of the experiment boric acid was completely bactericidal for group B streptococci at 40 g/l and partially so for a few strains at 20 g/l. Effective bacteriostasis was noted at 10 g/l and was incomplete at 2.5 g/l. Group D streptococci survived well in higher concentrations. Boric acid was fungistatic for C albicans. S saprophyticus was underrepresented in our experiment, as younger women with cystitis rarely present in hospital. The strains that were tested came from the hospital's staff clinic.

It emerges that for nearly all the bacteria and fungi found in urine boric acid is purely bacteriostatic at a range of concentrations between 10 g/l and 20 g/l. In our hands a few strains of group B streptococci showed a limited bactericidal effect at the concentration generally used and higher concentrations were lethal for all of them. For one of 17 of the strains of Ps aeruginosa, and for perhaps six of 14 of the A calcoaceticus strains tested 10 g/l was mildly bactericidal, though for many bacteriostasis was re-established at 20 g/l.

The results of the third experiment throw light on the practical importance of these indications of toxicity, which in any case involve only a proportion of some less common urinary pathogens. A calcoaceticus is not affected until it has been exposed to boric acid at a concentration of 10 g/l for over four hours. At the concentration normally used the small reduction in count seen at 24 hours was not evident at six hours. This organism is usually a hospital pathogen,14 so specimens containing it should not be subject to long delay before they are examined. With Ps aeruginosa the toxicity observed was restricted to concentrations of boric acid that are lower than usual. From graph h (figure) it can be seen that boric acid was toxic for some group B streptococci at 20 g/l. When one of these strains was tested in detail the marginal reduction in count was not apparent until the second hour of exposure and it did not then progress. The bactericidal effect was seen at 10 g/l. These observations show that boric acid is rarely toxic, and suggest that any limited toxicity has little practical relevance because it can often be avoided.

Our results show that although bacteria differ in their sensitivity to it, boric acid at a concentration of 18 g/l or 20 g/l is a sound choice for the preservation of urine. They also show that in certain circumstances there might be an advantage in using it at 10 g/l. Unless boric acid is emptied from a collection bottle, overdilution is impossible if the container is designed to be filled nearly to the top. Experience shows that bottles are often incompletely filled, particularly when specimens come from infants, so underdilution is a greater problem. If a bottle is less than half filled a bacterial count of 105/ml would be reached. This danger would be minimised by halving the concentration of boric acid, adding 0.25 g instead of 0.5 g to a 25 ml container. Against this must be set the incomplete and delayed paradoxical toxicity of boric acid at 10 g/l for a small proportion of some less common urinary pathogens. Alternatively, a smaller bottle containing an appropriately reduced amount of boric acid could be supplied, at least for specimens from infants.

Another advantage of the use of boric acid is that the lower numerical watershed can be adopted to separate positive from negative bacteriological reports. Practice already varies between laboratories in the choice of bacterial counts of 104/ml or 105/ml to divide negative from positive, and those who give an equivocal report on specimens with counts between these figures.15 There is evidence that preservation with boric acid clarifies the meaning of equivocal (104–105/ml) counts, making it more likely that they represent true infections.3 We suggest that boric acid deserves serious consideration as a general addition to specimen bottles used to transmit urine to bacteriological examination. Toxicity is rare and of theoretical rather than practical concern. The advantages of preservation with boric acid seem to outweigh greatly any disadvantages.

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