Sensitivity and rapidity of blood culture bottles in the detection of cornea organ culture media contamination by bacteria and fungi

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Aims: To test the bactericidal activity of standard organ culture medium, and to compare the sensitivity and rapidity of blood culture bottles with conventional microbiological methods for detection of bacteria and fungi inoculated in a standard cornea organ culture medium.

Methods: The bactericidal activity of contaminated standard organ culture medium containing 100 IU/ml penicillin, 0.1 mg/ml streptomycin, and 0.25 ug/ml amphotericin B was evaluated after 48 hours of incubation at 31°C with five inocula of 14 bacteria. Two yeasts (Candida spp) and one Aspergillus were also tested. Contaminated media were then inoculated in three blood bottles (aerobic, anaerobic, fungal) placed in a Bactec 9240 automat; three conventional microbiological broths were the control. Changes in colour of organ culture medium and growth on conventional broth were screened daily by visual inspection. The sensitivity and rapidity of detection of contamination were compared between the three methods: blood bottle, conventional, and visual.

Results: Orga culture medium eradicated five bacteria irrespective of the starting inoculums: Staphylococcus pneumoniae, Branhamella catarrhalis, Escherichia coli, Propionibacterium acnes, and Haemophilus influenzae. For micro-organisms where the medium was ineffective or bactericidal only (methicillin resistant Staphylococcus aureus, methicillin sensitive Staphylococcus aureus, Staphylococcus epidermidis, Staphylococcus haemolyticus, Pseudomonas aeruginosa, Acinetobacter baumannii, Bacillus subtilis, Klebsiella pneumoniae, Enterococcus faecalis, Candida albicans, Candida kruzei, Aspergillus fumigatus), the blood bottle, conventional, and visual methods detected microbial growth in 100%, 76.5%, and 70% of cases respectively. Mean detection time using blood bottles was 15.1 hours (SD 13.8, range 2–52). In cases of detection by the blood bottle method and the conventional method, the former was always faster: 95.5% against 65.2% detection within 24 hours (p=0.022) respectively.

Conclusions: Blood bottles detect more efficiently and more rapidly a wider range of bacteria and fungi than the conventional microbiological method and the visual inspection of organ culture media.
MATERIALS AND METHODS
The design of our experiments is presented in Figure 1.

Micro-organisms
Bacterial and fungal strains were obtained from the American Type Culture Collection (ATCC) (Rockville, MD, USA) or from corneal swabs and scrapings submitted for culture to the microbiology department of our hospital. Fourteen bacteria were tested: methicillin sensitive Staphylococcus aureus ATCC 25923, one methicillin resistant Staphylococcus aureus, Staphylococcus epidermidis ATCC 12228, one Staphylococcus haemolyticus, Streptococcus pneumoniae ATCC 49619, Enterococcus faecalis ATCC 29212, Haemophilus influenzae ATCC 49247, Branhamella catarrhalis ATCC 2631, Escherichia coli ATCC 25922, Klebsiella pneumonias ATCC 104, Pseudomonas aeruginosa ATCC 27853, Acinetobacter baumannii ATCC 19606, Bacillus subtilis ATCC 6633, and one Propionibacterium acnes. Two yeasts and one filamentous fungus were tested: Candida albicans ATCC 90028, Candida krusei ATCC 6258, and Aspergillus fumigatus ATCC 13935.

Bactericidal activity of the organ culture medium
The bactericidal activity of a commercial medium (Inosol, Chauvin-Opsia, Toulouse, France) containing 100 IU/ml penicillin G, 0.1 mg/ml streptomycin and 0.25 µg/ml amphotericin B, was measured on four inocula of the 17 micro-organisms. An overnight culture in Mueller-Hinton broth was briefly diluted in the organ culture medium to give $10^7$, $10^5$, $10^3$, and $10^2$ colony forming unit/ml (cfu/ml). A 100 µl aliquot was removed, serial diluted, and plated on Mueller-Hinton agar to determine the exact starting concentration of bacteria and fungi. The same amount of inoculum was added to Mueller-Hinton broth as a growth control. For the 14 bacteria, 0.1 ml of the organ culture medium incubated at 31°C was then removed at fixed times (2, 6, 24, and 48 hours) and plated on Mueller-Hinton agar (except for Streptococcus pneumoniae, Branhamella catarrhalis, and Propionibacterium acnes, where chocolate media were used). The number of colonies was counted after 48 hours’ incubation at 35°C and the results plotted on semilogarithmic graph paper. A bactericidal effect, defined as a $4 \log_{10}$ decrease in cfu/ml, corresponding to survival of less than 0.01% of the micro-organisms, was measured for the $10^7$ and $10^5$ inocula. Eradication was defined as a lack of surviving micro-organisms in the $10^3$ and $10^2$ cfu/ml inocula. The $10^2$ cfu/ml inoculum was chosen to assess how well the three methods detect very low contamination. Each inoculum for each micro-organism was tested three times. Given the very low concentration of amphotericin B in the organ culture medium, fungicidal activity was not tested.

The inoculated organ culture media were incubated in sealed flasks for 48 hours at 31°C in a conventional carbon dioxide free dry incubator, identical to that used in our cornea bank. In most European banks’ corneas routinely spend this period in the so called “transport” medium before the first endothelial and microbiological tests are done and the medium changed.
Sensitivity and rapidity of the three methods in detecting contamination

In the visual method, changes in colour (to orange or yellow) or in turbidity of the organ culture medium, indicating positivity, were screened daily for 14 days by visual inspection.

In the conventional method, 1 ml of contaminated organ culture medium was inoculated in a brain-heart broth (10 ml) and a Schaedler broth (10 ml) incubated at 35°C and in one Sabouraud agar (10 ml) incubated at 28°C. Growth on conventional broth was screened daily by visual inspection for 14 days.

In the blood bottle method, 2.5 ml of contaminated organ culture medium was injected into one Bactec Plus Aerobic/F bottle (25 ml), one Bactec Lytic/10 Anaerobic/F bottle (40 ml), and one Bactec Mycosis IC/F (40 ml) (Becton Dickinson, Cockeysville, MD, USA). The bottles were placed in a Bactec 9240 incubator for 14 days at 35°C and rocked continuously. As described in a previous paper, the incubator detected any rise in carbon dioxide produced by bacterial growth. A sensor placed at the bottom of each bottle reacted with the carbon dioxide and produced fluorescence proportional to the carbon dioxide level. Fluorescence was measured every 10 minutes and time to detection was rounded to the nearest hour.

In all three methods, identification of isolate micro-organism was verified from subcultures of the positive media by standard bacteriological methods. The inoculated micro-organism always corresponded to the one detected, thus ruling out any exogenous contamination during manipulation by the technician.

RESULTS

Bactericidal activity of the organ culture medium (Table 1)

The antibiotics in the organ culture medium eradicated five bacteria irrespective of the starting inoculum: Streptococcus pneumoniae, Branhamella catarrhalis, Escherichia coli, Propionibacterium acnes, and Haemophilus influenzae. The antibiotics of the organ culture medium had no effect on the methicillin-resistant Staphylococcus aureus. Only the 10⁷, 10⁸, and 10⁹ inocula of Staphylococcus haemolyticus and Bacillus subtilis were eradicated.

Sensitivity and rapidity of the three methods in detecting residual contamination (Table 2)

Detection of residual contamination was 100% by the blood bottles (30/30), 76.5% by the conventional method (23/30), and only 70% by the visual method (21/30). No blood bottle tested positive without an identifiable germ, thus ruling out the possibility of a false positive having been generated by a physicochemical reaction of the organ culture medium with the contents of the bottle. When both the conventional and the blood bottle methods detected contamination (n=23), the latter was significantly quicker: positivity was nearly always observed within 24 hours (95.5%, 22/23 against 65.2%, 15/23) (Fisher exact test, p=0.022). Blood bottles also detected contamination within 8 hours in 40% of cases (12/30), corresponding to the technicians’ shift length. Mean time to contamination detection by the blood bottles was 15.1 hours (SD 13.8, range 2–52): 19.5 hours for the bacteria (SD 14.2, range 3–52), and only 3 hours (SD 0.4, range 2–4) for the yeasts.

DISCUSSION

Early post-PKP endophthalmitis may be linked to the recurrence of an infection present in the recipient before grafting, to peroperative contamination, or to transmission of a micro-organism via contaminated graft tissue. Organ culture at 31°C, 34°C, or 37°C, because of the presence of antibiotics at these temperatures, coloured pH indicators and microbiological tests, eradicates micro-organisms or at least detects residual survival. It thus made sense, in the first series of experiments, to test the bactericidal strength of the organ culture medium. We used an original microbiological protocol that adapts techniques for studying the kinetics of antibiotic activity (time killing curves) to the real cornea storage temperature (31°C in our bank). The second series of experiments compared how effectively and quickly residual contamination is detected by three methods: a bacteriological method thought more suitable for low inocula and/or when antibiotics are present, a conventional microbiological method, and daily visual inspection of the colour of organ culture media.

The micro-organisms studied were chosen because they are implicated in early post-PKP endophthalmitis or because they are typically present in postmortem eye flora. The combinations of antibiotics and antifungals used in organ culture media in Europe are similar. The 2002 survey of 78 cornea banks by the European Eye Bank Association showed that 68% (53/78) used organ culture alone or combined with storage at 4°C. Penicillin was combined with streptomycin in 87% of cases (66/73), nearly always in concentrations of 100 IU/ml and 0.1 mg/ml respectively. According to the same survey, an antifungal is always used: amphotericin B is most common (92.5%, 49/53

| Table 1 Bactericidal activity and eradication capacity of organ culture medium according to the bacteria and the inoculum. Because of the definition of bactericidal activity, * for 10⁷ and 10⁸ inocula, only eradication was checked. |
|---|---|---|---|---|---|---|
| Inoculum | 10⁷ | 10⁸ | 10⁹ | 10⁷ | 10⁸ | 10⁹ |
| **Bactericidal** | **Eradication** | **Eradication** | **Bactericidal** | **Eradication** | **Bactericidal** | **Eradication** |
| Methicillin resistant Staphylococcus aureus | no | no | no | no | no | no |
| Methicillin sensitive Staphylococcus aureus | yes | no | yes | no | yes | yes |
| Staphylococcus epidermidis | no | no | no | no | no | no |
| Staphylococcus haemolyticus | yes | no | yes | yes | yes | yes |
| Pseudomonas aeruginosa | no | no | no | no | no | no |
| Acinetobacter baumannii | no | no | no | no | no | no |
| Bacillus subtilis | no | no | yes | yes | yes | yes |
| Klebsiella pneumoniae | no | no | yes | no | yes | yes |
| Enterococcus faecalis | yes | no | yes | no | yes | yes |
| Streptococcus pneumoniae | yes | yes | yes | yes | yes | yes |
| Branhamella catarrhalis | yes | yes | yes | yes | yes | yes |
| Escherichia coli | yes | yes | yes | yes | yes | yes |
| Propionibacterium acnes | yes | yes | yes | yes | yes | yes |
| Haemophilus influenzae | yes | yes | yes | yes | yes | yes |

* Bactericidal activity = 4 log₁₀ decrease in colony forming units/ml after 48 hours.
† Eradication = lack of survival of bacteria after 48 hours.
cases) although concentrations differ between a trace (0.25 µg/ml) in 77.5% of cases (38/49), low (2.5 µg/ml) in 18.5% of cases (9/49), and high (25 µg/ml) in two banks (4%). The combination of penicillin and streptomycin was thus chosen in our experiment because they are very common. This combination of penicillin and streptomycin was thus chosen in our experiment because they are very common. This combination was highly effective, eradicating five bacteria. This result agrees with previous studies of similar organ culture media. However, its transmission via the grafted cornea was proved in only two cases; more importantly, the corneas were stored at 4°C in a medium not containing a penicillin active on this organism.

Similarly, Propionibacterium acnes and Streptococcus pneumoniae have been found in corneoscleral rims, but only after storage at 4°C. In that review, how-
The superiority of blood bottles appears clearly. We showed them to be more sensitive than the conventional microbiological method with the visual method. They detected a wider spectrum of bacteria, sometimes even with high inocula missed by the other two techniques. Methicillin sensitive <i>Staphylococcus aureus</i> at 10<sup>3</sup> and 10<sup>4</sup> cfu/ml, <i>Staphylococcus haemolyticus</i> at 10<sup>5</sup> cfu/ml, <i>Klebsiella pneumoniae</i> at 10<sup>2</sup> cfu/ml, and <i>Enteroococcus faecalis</i> at 10<sup>1</sup> cfu/ml were never detected by the conventional method and induced no change of colour of the organ culture medium. Hughes demonstrated that culture using blood bottle methods detected significantly more pathogens than conventional methods, especially of <i>Staphylococcus</i> and <i>Enterococcus</i>. This is important in ophthalmology because these are among the main organisms responsible for post-PKP endophthalmitis.<sup>31</sup>

In cases of low inocula, the high sensitivity of blood bottles was predictable. The lack of sensitivity of the conventional microbiological method may be due to antibiotics contained in the organ culture media and which may inhibit the growth of the micro-organism in culture. Resins, such as those present in Bactec Plus Aerobic/F and Bactec Lytic/10 Anaerobic/F bottles, may reduce the inhibiting effects of antibiotics.<sup>25</sup> It is likely that these low inocula would have been eradicated when the medium was changed: fresh antibiotics are then added, an important point because their action is limited in time.<sup>35</sup>

Lastly, it is likely that a very low inoculum has no contaminating potential in the recipient because of the bac-tericidal activity of the aqueous humour and because of the perioperative antibiotics systematically prescribed. Highly sensitive detection methods, such as Alborn's, subjecting corneoscleral rims to mechanical extraction in a Stomacher blender, detected micro-organisms in 28.9% of cases, whereas microbiological testing of organ culture media found them in only 0.53%<sup>26</sup> to 0.7%<sup>27</sup> of cases. Yet Alborn, in her series, reported no endophthalmitis. In our study, low inocula of 10<sup>5</sup> and 10<sup>6</sup> cfu/ml of methicillin resistant <i>Staphylococcus aureus</i> were detected only by blood bottles, in 36 and 39.3 hours respectively, and would probably have been eradicated at the renewal of organ culture medium. We think that, in this respect, the lowering of the sensitivity threshold is more useful for quality control of the entire storage procedure, from procurement to graft, than to surgeons as a predictor of post-operative infection.

Given the very low concentration and the instability of amphotericin B, which is used mainly to ensure fungicidal action during production of the organ culture medium, we felt it unnecessary to test the fungicidal activity of the organ culture medium. We were reinforced in this choice since the three fungi grew and were detected after the 48 hour incubation in the organ culture medium irrespective of the inoculum. The rarity yet severity of fungal endophthalmitis,<sup>13</sup> 24 and the slowness of growth for some species, naturally led us to optimise the microbiological method may be due to antibiotics contained in the blood bottles and conventional microbiological media only once daily. This consolidates the utility of the blood bottles, which detected contamination within 8 hours in nearly half the cases, corresponding to the technicians’ shift length. Early elimination from storage of contaminated corneas, particularly due to yeasts, minimises the risk of contamination of the other corneas in the incubator and limits storage costs. In addition, replacement of the conventional method with an automated culture system like the Bactec blood bottle method significantly reduced labour requirements in our bacteriology laboratory; and, despite the extra cost of the three blood bottles (approximately €10 + €1.9; €6.3 + €1.2), yielded substantial savings. Lastly, our use of this protocol at the start and end of organ culture allows faster delivery of corneas without affecting their microbiological safety, and our patients benefit from corneas of higher endothelial cell density.

CONCLUSION

Use of this more sensitive protocol should, in theory, cause a higher cornea discard rate. But in practice it is likely that, because there are fewer operation steps performed in better conditions (blood bottles are sealed, and specimens can be directly injected under laminar flow hood in the cornea banks or operating theatre), this effect is offset by a fall in exogenous contamination. The contamination rate of 3.2% (30/934 corneas), obtained by this protocol since its introduction in our cornea bank and by the favourable clinical outcome in our patients (one hypopyon out of 412 penetrating keratoplasty), is most acceptable.

We consider that the greater sensitivity and speed, at no extra cost, of blood bottles in testing the microbiological safety of organ culture media compared to conventional microbiological methods, justifies their general introduction for routine use.

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