Microbial decontamination of human donor eyes with povidone-iodine: penetration, toxicity, and effectiveness

E Pels, G F J M Vrensen

Abstract

Background/aims—Povidone-iodine (PVP-I) is applied for microbial decontamination of human eyes donated for transplantation. Concentrations and immersion times vary greatly. The effectiveness and toxicity of PVP-I were assessed for different decontamination protocols.

Methods—Human donor eyes and corneas were immersed in different concentrations (5–100 mg/ml) of PVP-I for different times (2–30 minutes). The penetration of iodine into the corneal tissue was assessed by x-ray microanalysis. Microbial contamination was determined by taking cultures of the limbal areas and storage solutions and by incubation of the corneoscleral buttons in antibiotic-free culture medium. Cytotoxicity of PVP-I for corneal fibroblasts in culture was assessed using the MTT assay.

Results—Depending on concentration and immersion time iodine was found to penetrate into the epithelium, Bowman’s layer, and stroma in amounts equivalent to 2–40 mg/ml PVP-I. The MTT assay demonstrated that 2.5 mg/ml PVP-I caused total damage to fibroblasts in vitro. Rinsing eyes with tap water and subsequent immersion in PVP-I reduced the rate of contamination from 82 out of 106 to 69 out of 106 and 37 out of 106, respectively. Antibiotics in the storage medium further reduced contamination from about 40% to 3%. Microbial contamination was not reduced by increasing the concentration and immersion times beyond 5 mg/ml PVP-I for 2 minutes.

Conclusion—Immersion of human donor eyes in 5 mg/ml PVP-I solution for 2 minutes significantly reduces microbial contamination of donor corneas without relevant penetration of iodine into the corneal layers. Higher PVP-I concentrations and longer immersion times do not further reduce contamination, whereas the amount of iodine penetrating the corneal layers is elevated above the level cytotoxic for corneal fibroblasts. In view of this, concentrations above 5 mg/ml of PVP-I and immersion periods over 2 minutes are not recommended for reduction of the contamination rate of donor eyes.

All cornea banks have to cope with the problem of supplying viable donor tissue free from microbial contaminants, while donor eyes are generally contaminated. Different decontamination procedures have been advocated: (i) treatment with antibiotics; (ii) irrigation; (iii) mechanical abrasion of the epithelium; and (iv) immersion of eyes in povidone-iodine/betadine. The use of povidone-iodine (PVP-I, polyvinylpyrrolidone-iodine complex) is attractive because of its broad antimicrobial action spectrum including most Gram positive and Gram negative bacteria, fungi, yeasts, viruses, and protozoa. In addition, it is chemically stable, inexpensive, and readily available. For the same reasons PVP-I is suggested as treatment for corneal ulcers in cases when antibiotics are not available as, for example, in developing countries.

The concentration of “free” iodine significantly contributes to the antimicrobial activity of PVP-I. The effect of hypiodous acid which also contributes to the antimicrobial activity of PVP-I is not considered in this paper. Dilution of PVP-I results in weakening of the iodine binding to the carrier molecule and a concomitant increase in the amount of “free” iodine in the solution. In antiseptic practice watery solutions of 50–100 mg/ml PVP-I are commonly used. However, for decontamination of human donor eyes concentrations of 5–100 mg/ml and immersion times of 2–10 minutes are applied by eye banks. Despite the reported low ocular toxicity of PVP-I some toxic effects have been described for the corneal epithelium and fibroblasts at concentrations of 25 and 50 mg/ml and immersion periods over 4 minutes. Recently, it has been shown that 50 mg/ml PVP-I for 3 minutes is superior to ciprofloxacin and gentamicin in antimicrobial effectiveness, but corneal tolerance of the procedure still needs investigation.

In this study energy dispersive x-ray microanalysis (EDXM) was performed to investigate the penetration of iodine into the cornea and its dependence on PVP-I dose, immersion time, and intactness of the epithelial layer. In addition, the concentration dependent toxicity of PVP-I for human corneal fibroblasts was assessed using the MTT assay and was related to the amount of iodine present in the cornea. The effectiveness of different concentrations of PVP-I and immersion times for bacterial and fungal decontamination of corneoscleral buttons was evaluated.

Methods

HUMAN CORNEAS

Human eyes were donated to the Foundation BIS (Leiden, Netherlands) for transplantation.
purposes. Eyes were enucleated by certified enucleators using aseptic techniques. Antibiotics were not installed in the eye and copious irrigation was not carried out before enucleation. Transportation of the eyes to the cornea bank by BIS took place in a moist chamber on ice. No antibiotics were present in the bathing saline solution. Corneas judged unsuitable for transplantation because of small scars in the optic centre or senile changes in the endothelium were used for the present experiments.

POVIDONE-IODINE

Solutions were prepared from powdered polyvinylpyrrolidone-iodine (PVP-I, Sigma Chemicals Co, St Louis, MO, USA). For the penetration and decontamination experiments PVP-I was dissolved in deionized water (Milli Ro4 and Q Reagent Water systems, Millipore, Bedford, MA, USA) in concentrations of 5, 20, 50, and 100 mg/ml and pH was adjusted to 6.0 with 1 N NaOH. Solutions were filtered (0.22 µm pore size, Stericup GS, Millipore, Bedford, MA, USA), aseptically divided (30 ml/vial) in sterile wide necked 50 ml vials (Greiner GmbH, Frickenhausen, Germany), and stored in the dark at room temperature. For toxicity experiments a 10 mg/ml PVP-I solution in sterile phosphate buffered saline (PBS), (Azur Pharmacy, Amsterdam, Netherlands) was prepared, filtered (0.22 µm pore size), and aseptically diluted with sterile PBS shortly before use.

X Ray Microanalysis

Procedure

Corneal tissue (see below) was snap frozen in liquid nitrogen, vacuum freeze dried at −0°C (Virtis Benchtop 5, Virtis Co, Gardiner, NY, USA) and stored on silica gel. The bulk specimens were mounted on a carbon holder and left uncoated. Inspection and analysis was carried out in a Philips SEM 505 scanning electron microscope (Philips Industries, Eindhoven, Netherlands) equipped with an Edax PV 9800 microanalysis system using a windowless ECON detector (Edax PB 9760/26, Edax Ltd, Mahwah, NJ, USA). A high tension of 15 kV was used throughout and spectra were obtained from spot measurements (spot size 200 nm) at the transverse sides of cut corneal pieces in (i) the basal layer of the epithelium, (ii) Bowman’s layer, (iii) anterior stroma, and (iv) mid-stroma (arrowheads, Fig 1). The peak to background ratio (P/B ratio) was used to determine the local mass of iodine. The P/B ratio provides an intrinsic absorption correction independent of surface geometry and therefore suitable for analysis of biological bulk specimens. Specimens were randomly numbered for bias-free monitoring of the presence of iodine. Owing to the dehydrating effect of povidone, however, corneal samples treated with higher concentrations or increased immersion times of PVP-I were nevertheless easily recognised because of the reduced thickness of the cornea.

Tissue selection

Reliable EDXM quantification of elements in biological materials strongly depends on keeping all variables constant including those of the tissue investigated. Therefore we decided not to use whole corneas for the different concentrations of PVP-I and different immersion times to be tested but to divide excised corneoscleral buttons into six triangular pieces. In order to see whether the use of pieces reflects the actual situation of rinsing the whole bulb we carried out the following pilot experiment. From paired eyes one globe was immersed in 50 mg/ml PVP-I for 5 minutes before excision of the corneoscleral button while from the fellow eye the corneoscleral button was excised, divided into six triangular pieces each, and subsequently immersed in 50 mg/ml PVP-I for 5 minutes. A control eye and control pieces were immersed in PBS. X Ray spectra were taken from the pieces at the epithelial, anterior stroma, and mid-stroma sites and at comparable sites of the intact corneas. The X ray spectra and P/B ratios of I were nearly identical at these sites in the pieces compared with those in the intact cornea. Although iodine concentrations at the lateral edges and posterior stroma were higher than in the intact cornea under the
specific conditions (given the penetration of PVP-I from the lateral and posterior sides of the pieces) they did not influence the concentration in the middle of the pieces. However, to avoid even the smallest influence on PVP-I measurement from PVP-I present at the edges, 1 mm of tissue was removed from the lateral sides with a razor blade.

Another relevant practical point is whether the presence or absence of the epithelium affects the penetration of PVP-I into the cornea. To test this in one of two fellow eyes the epithelium was removed by scraping with a razor blade. The other eye was used as control. Both bulbs were subsequently immersed in various concentrations PVP-I for various times and treated as described above.

Calibration of EDX iodine signals
Contrary to the situation in material sciences, where for quantitation it is mostly sufficient to know the ratio in which the different elements are present in the sample, biological specimens require calibration standards for quantititation preferably resembling the specimen in chemical and physical properties. In this study standard specimens were prepared from 15% collagen I from rat tails, equivalent to the amount of collagen in the corneal stroma and 5% agarose type VII (Sigma Chemical Co, St Louis, MO, USA) with a low gelling temperature to substitute for the 5% non-collagen protein in the stroma and to give the standard specimens some matrix structure. Because corneal stroma may vary locally in protein content at different hydration states, low protein standards with one fifth of the collagen concentration were also prepared. PVP-I was added to this mixture in concentrations of 0–100 mg/ml before gelling. In three independent experiments a highly significant linear correlation was observed between the PVP-I concentration in the standard specimens and the P/B ratios of iodine. The protein content of the standard specimens did not affect this correlation (Fig 2). The detection limit lies at about 2 mg/ml PVP-I which corresponds to 0.2 mg/ml iodine.

MTT ASSAY TO DETECT PVP-I TOXICITY FOR CORNIAL FIBROBLASTS

Tissue
Corneal fibroblasts were obtained from normal human donor eyes as described. In short, corneal buttons were excised and cut in small pieces. Three to four corneal pieces were implanted per well of a six well plate (Nunc, Roskilde, Denmark) and incubated with 1 ml Eagle’s modified minimum essential medium (EMEM, ICN Biomedicals Inc, Costa Mesa, CA, USA) supplemented with 10% fetal bovine serum (FBS, Gibco Life Technologies Ltd, Paisley, Scotland), 100 units/ml penicillin (Gist Brocades, Leiderdorp, Netherlands), and 50 μg/ml streptomycin (Biochemie GmbH, Vienna, Austria) at 37°C. When the pieces adhered firmly to the bottom of the well by the outgrowing cells 5 ml medium per well was added. Medium was renewed twice a week. Outgrowing fibroblasts were subcultured after 25–28 days and fibroblasts of the second passage were stored in liquid nitrogen. Cells were defrosted, subcultured for one passage, and used for experiments. At confluency they were removed from the culture flasks by incubation with a trypsin/EDTA solution for 5 minutes, collected by centrifugation (200 × g), resuspended in medium, and plated out at 5 ×10⁴ cells per well in 96 well flat bottomed tissue culture plates (Nunc, Roskilde, Denmark). The cells were cultured at 37°C in a humidified atmosphere with 5% carbon dioxide before exposure to freshly prepared PVP-I solutions, varying in concentration from 0.01 to 10 mg/ml PVP-I.

Procedure
Cytotoxicity was determined with the MTT assay directly and 48 hours after exposure to PVP-I. The MTT assay basically measures the capacity of mitochondrial enzymes to transform the MTT (3-(4,5 dimethyl thiazole-2yl)-2.5 diphenyl tetrazolium bromide; Sigma Chemicals Co, St Louis, MO, USA) salt into a formazan product and thus reflects the mitochondrial activity of corneal fibroblasts. After exposure of the cells to the PVP-I solutions (2 minutes, 50 μl/well), the cells were carefully washed three times with PBS and 100 μl of a culture medium with 2% FBS was added per well. Directly (acute test) or after 48 hours (delayed test) 10 μl of a freshly prepared MTT solution (5 mg/ml MTT in PBS) was added to all wells and plates were incubated at 37°C for 4 hours. The MTT formazan reaction product was solubilised by the addition of 200 μl acid-isopropanol (0.04 N HCl in propanol-2) to the incubation medium without washing steps and overnight incubated at room temperature in the dark. Optical density of the solution was assessed at 560 nm in a spectrophotometer (I EMS Reader MF, Labsystems, Helsinki, Finland) taking medium with MTT and isopropanol as the blank. Each experiment was performed in eightfold. The experiments were repeated five times. The damage was calcu-
CONTROL OF ASEPTIC PROCEDURES USED

To exclude the possibility that the observed contamination of corneas was induced by improper aseptic techniques and working surroundings, working conditions were evaluated. Corneas \((n=250)\) routinely stored for 14–24 days and judged unsuitable for transplantation after storage were subsequently processed including light microscopic examination and stored in medium without antibiotics at 31°C for another 4 weeks or until contamination become obvious. Tissue contamination should have revealed itself within the first 14–24 days of storage, while improper working conditions would become manifest in the subsequent storage period in the medium without antibiotics.

EFFECTIVENESS OF PVP-I SOLUTIONS

Donor eyes with corneas unsuitable for transplantation because of scars in the optic centre as observed by slit lamp examination or because of senile changes of the endothelium as observed in the fellow cornea, were used for this part of the study. After removal of adnexal and muscle tissue from the globe using sterile instruments, eyes were rinsed with tap water for 1 minute. After rinsing, the eyes were immersed in different PVP-I solutions (5, 20, 50, and 100 mg/ml) for 2 or 10 minutes, subsequently in 0.5% sodium thiosulphate solution in PBS for 1 minute and finally rinsed with PBS. After preparation the corneoscleral button was dissected in two pieces. One was incubated in a sterile vial with storage medium as used for routine eye bank storage. The other was incubated in storage medium without antibiotics. Both were stored at 31°C for 4 weeks or until contamination became obvious. The groups consisted of 25–30 corneas each.

STATISTICS

Multiple regression between P/B ratios of iodine and PVP-I concentration and immersion time was determined giving regression coefficients and p values for the slope. Differences in P/B ratios in corneas with and without epithelium were compared using the Wilcoxon signed rank sum test. The proportion of contaminated corneas in the different treatment groups were compared using the \(\chi^2\) test. Level of significance was set at 5%.

Figure 3 EDAX spectra of a cornea immersed in 50 mg/ml PVP-I for 10 minutes analysed in the basal layer of the epithelium (A) and anterior stroma (B). For details see text.
Results

IODINE CONTENT IN CORNEAS ASSAYED BY X RAY MICROANALYSIS

The x ray spectra as exemplified in Figure 3, taken from PVP-I immersed corneas, clearly show the Lα and Lβ peaks of iodine. Moreover, the Kα peaks of C, O, P, and S are characteristic for biological material when using a windowless detector and indicate the presence of proteins, phospholipids, DNA, and RNA. In line with the higher density of membranes, nuclei, and ribosomes in the epithelium and the higher protein (collagen) content of the stroma the P peak is more pronounced in the epithelium (Fig 3A) and the S peak in the stroma (Fig 3B). There is no reasonable explanation for the high Kα peak of Cl also found in untreated material. This must be an artefact although, at present, of unknown origin.

The P/B ratios of I Lα at different corneal sites, epithelium, Bowman’s layer, anterior stroma, mid-stroma (see Fig 1), correlated significantly with PVP-I concentration and immersion periods (Fig 4A–D respectively). At the lowest concentration, iodine is observed in the epithelium at 5 minutes (Fig 4A) and at 30 minutes some iodine is found in the stroma (Fig 4C, D). After 2 minutes’ exposure of the higher concentrations (20, 50, and 100 mg/ml) iodine is observed in Bowman’s layer. Penetration of iodine into the stromal layers already occurs after 5 minutes’ exposure to the highest concentrations (50 and 100 mg/ml).

Table 1 Effect of epithelial removal on the P/B iodine ratio

<table>
<thead>
<tr>
<th>Concentration PVP-I solution*</th>
<th>20 mg/ml</th>
<th>100 mg/ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exposition time†</td>
<td>Corneal layer</td>
<td>P/B ratio iodine (mean (SD))</td>
</tr>
<tr>
<td>2 minutes</td>
<td>Epithelium</td>
<td>11 (2)</td>
</tr>
<tr>
<td></td>
<td>Bowman's layer</td>
<td>6 (4)</td>
</tr>
<tr>
<td></td>
<td>Anterior stroma</td>
<td>2 (0)</td>
</tr>
<tr>
<td></td>
<td>Mid-stroma</td>
<td>0</td>
</tr>
<tr>
<td>10 minutes</td>
<td>Epithelium</td>
<td>46 (13)</td>
</tr>
<tr>
<td></td>
<td>Bowman's layer</td>
<td>23 (7)</td>
</tr>
<tr>
<td></td>
<td>Anterior stroma</td>
<td>15 (4)</td>
</tr>
<tr>
<td></td>
<td>Mid-stroma</td>
<td>12 (5)</td>
</tr>
<tr>
<td>30 minutes</td>
<td>Epithelium</td>
<td>37 (11)</td>
</tr>
<tr>
<td></td>
<td>Bowman's layer</td>
<td>19 (7)</td>
</tr>
<tr>
<td></td>
<td>Anterior stroma</td>
<td>15 (9)</td>
</tr>
<tr>
<td></td>
<td>Mid-stroma</td>
<td>9 (8)</td>
</tr>
</tbody>
</table>

Signed rank test p <0.05 p >0.05

*Data other concentrations (10 and 50 mg/ml) not shown.
†Data other exposure times (5 and 15 minutes) not shown.
TOXICITY OF PVP-I FOR CORNEAL FIBROBLASTS

Significant damage of the corneal fibroblasts was observed at a concentration of 0.25 mg/ml PVP-I for 2 minutes (Fig 5). Lower concentrations gave no significant damage, neither directly nor 48 hours after immersion. Cell damage increased with increasing PVP-I concentrations. Nearly total damage was observed at concentrations of 2.5 mg/ml PVP-I and more both in the direct and in the delayed test. Substance concentrations at half maximum cytotoxicity (LC50) calculated from the dose-response curves, were 0.4 and 0.6 mg/ml for the direct and delayed test respectively (Sigmoidal fit, Fig 5).

MONITORING THE EFFECTIVENESS OF DECONTAMINATION PROCEDURES USED IN ROUTINE EYE BANKING

On arrival 82 of the 106 eyes were found to be contaminated with surface contaminants (Table 2). After rinsing with tap water the ratio of contaminated eyes decreased to 69 out of 106, while the number of colonies observed also decreased. Most cultures were pure consisting of one microbial strain. After treatment with 0.5% PVP-I for 2 minutes the ratio of contaminated eyes decreased further to 37 out of 106 (Table 2).

From the corneal pieces stored without and with antibiotics 34 out of 90 (38%) and four out of 90 (4%) turned out to be contaminated, respectively.

Table 2 Effect decontamination procedure on surface contaminants

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Contamination</th>
<th>Degree of contamination</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>no</td>
<td>&gt;20 colonies per plate</td>
</tr>
<tr>
<td>Control</td>
<td>24</td>
<td>82</td>
</tr>
<tr>
<td>After rinsing</td>
<td>37</td>
<td>69</td>
</tr>
<tr>
<td>After rinsing and PVP-I</td>
<td>69</td>
<td>37</td>
</tr>
<tr>
<td>Pearson χ²</td>
<td>41.87</td>
<td>10.85</td>
</tr>
<tr>
<td>Significance</td>
<td>0.0000</td>
<td>0.0044</td>
</tr>
</tbody>
</table>

MONITORING OF MICROBIAL CONTAMINATION DURING ROUTINE EYE BANKING

Control of tissue

Three per cent of corneas stored routinely by organ culture in the bank (1994–6 follow up study of 4000 corneas) turned out to be contaminated. Contaminants were Staphylococcus epidermidis (36%), Pseudomonas sp (27%), Escherichia coli (9%), and Bacillus sp (8%). Fungal contamination was less than 0.6%. Most contaminants (95%) were detected within a 3 day storage period. However, some escaped detection in the medium sample and showed up around day 10 in the storage solution or in the transport solution. No difference in contamination rate was observed between corneas accepted for transplantation or judged unsuitable.

Control of procedure

Corneas (n=250) were routinely processed, incubated for 14–24 days and removed from this medium for evaluation of the endothelium. They were returned to medium without antibiotics to discriminate whether contamination of corneas during storage is due to remaining contaminants or is induced by inadequate working conditions. Not one single cornea turned out to be contaminated.

EFFECTIVENESS OF PVP-I SOLUTIONS

Eyes were rinsed with tap water as usual and then submersed for 2 or 10 minutes in different concentrations of PVP-I (5, 20, 50, and 100 mg/ml). After excision the corneoscleral buttons were stored in medium without antibiotics. The effectiveness of the decontaminating procedure was expressed as percentage corneas observed free of contamination. No significant differences were observed between the different groups (Table 3). With a sample size of about 25 for each group a 20% difference between the treatment groups can be detected in 90% of trials. Staphylococcus epidermidis was again the most common contaminant (43%).

Discussion

After immersion of corneal tissue in PVP-I solutions as part of a decontamination process, iodine was found to penetrate the corneal tissue. Depending on concentration and exposure time iodine was found from the basal epithelium to half way in the stromal depth. At treatment with 5–50 mg/ml PVP-I for 2 minutes the iodine was virtually absent in the stromal solution or in the transport solution. No differences were observed between corneas accepted for transplantation or judged unsuitable.

Table 3 Effectiveness of PVP-I

<table>
<thead>
<tr>
<th>% corneas not contaminated (n/ ntotal)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exposure time (minutes)</td>
</tr>
<tr>
<td>2</td>
</tr>
<tr>
<td>10</td>
</tr>
<tr>
<td>PVP-I 5 mg/ml</td>
</tr>
<tr>
<td>PVP-I 20 mg/ml</td>
</tr>
<tr>
<td>PVP-I 50 mg/ml</td>
</tr>
<tr>
<td>PVP-I 100 mg/ml</td>
</tr>
</tbody>
</table>

Figure 5 Dose/response curve of PVP-I solutions for human corneal fibroblasts. Damage was assessed with the MTT assay directly after exposure of the cells to the PVP-I and 48 hours later. Mean (SD) of the means obtained in five experiments in eightfold. Asterisks indicate damage significantly different from controls.
methods for quantitative analysis of bulk specimens. In order to get an indication of the amount of iodine penetrated, standards were prepared resembling the composition of the corneal stroma. Because PVP-I exerts a dehydrating effect on the corneal stroma and postmortem corneas vary in hydration state, standards with high and low protein content were used. No differences were observed between the high and low protein standards provided the amount of iodine was expressed as the peak to background ratio. This P/B ratio was linearly correlated with the iodine content of the standards. The minimum amount of iodine detected corresponds to 0.2 mg/ml (2 mg/ml PVP-I), and this is far above the claimed levels of PVP-I were found for corneal reduction of enzyme activity. Similar toxicity of PVP-I is not a temporarily reversible test with 0.5 mg/ml PVP-I, indicating that the effect occurred in both the acute test and the delayed mal fibroblasts in vitro. Maximal damage PVP, or as iodide after chemical reduction. The amount in the stroma is low compared with that in the epithelium but it is apparently sufficient to cause fibroblast damage and death with immersion periods of more than 2 minutes and concentrations above 5 mg/ml PVP-I.

Because of its broad action spectrum and low toxicity for the epithelium PVP-I may be a useful treatment for corneal ulcers. In that case toxic effects on the fibroblasts, because of its penetration into the corneal tissue, are of minor importance. In routine eye banking decontamination of human donor eyes with 5 mg/ml PVP-I for 2 minutes significantly reduces the viability of microbial surface contaminants and thus the risk of corneal tissue contamination. However, it does not eliminate all contaminating microbes. After decontamination procedures 36% of the eyes remained positive in microbiological tests. Higher concentrations of PVP-I and longer immersion periods do not further reduce the percentage of contaminated corneas. These results are in agreement with studies demonstrating more rapid bactericidal action of diluted solutions of PVP-I than a full strength 10% solution. The free iodine, the reactive component of the PVP-I complex, is constantly released in small amounts, remains in equilibrium with the complex, and undergoes chemical reactions characteristic of iodine until the available iodine is exhausted. It follows a bell shaped curve with a maximum level in a 7 mg/ml solution. The bactericidal activity of PVP-I decreases by exposure to organic substances. Therefore, removal of excess tissue from the eye such as remnants of muscles, conjunctiva and orbital fat, is important for optimal results.

The rate of contamination among eye banks using organ culture varies from 0.53% to 5% and a mean percentage of 3.5% is reported for banks in Europe. The cause of death and the time elapsed from death to enucleation may have a marked influence on the risk of contamination. Improved removal of contaminants early in the process to minimise the contaminating microbial load and the amount of bacterial waste products such as endotoxins in the storage solution is of course advantageous.

Antibiotics in the storage solution were shown to be needed as a last step of the decon-
tamination procedure to reduce the microbial risk to an acceptable level. One approach is to accept a certain level of contamination with a conventional penicillin/streptomycin containing medium. If, in that case, an organism resistant to the antibiotics is present in low numbers and escapes detection during the sterility controls a wide range of more effective antibiotics is still available for the clinician in cases of phthisa. Another approach is to keep the contamination rate at nearly zero levels with wide spectrum and powerful antibiotics such as amoxicillin. However, in that case multiresistant microbes may be induced in organ culture surroundings favouring antibiotic resistance development. In case these microbes enter the eye after grafting, antibiotic treatment is not available.

After immersion PVP-I remains on the outside of the bulbus. With increasing concentrations the viscosity of the PVP-I solution increases and it becomes more difficult to remove the reactive iodine before preparation of the corneoscleral button. During excision of the corneoscleral button PVP-I may inadvertently penetrate the anterior chamber and may damage the endothelium. The sodium thiosulphate solution advocated to reduce the activity of the remaining free iodine is useful in these cases and might itself inhibit bacterial growth.

It is evident that the effectiveness of decontamination procedures can only be evaluated correctly if the working conditions in the bank, such as aseptic handling for storage and evaluation of the corneoscleral buttons, do not add to the risk of contamination. This was shown to be the case in our cornea bank.

The present study allows the following conclusions:

(i) Rinsing with tap water and immersion in PVP-I solution together with classic antibiotics in the storage solution reduce the contamination rate of human donor corneas to 3% under the conditions studied;

(ii) PVP-I (0.5%) 5 mg/ml for 2 minutes is sufficient for maximal decontaminating effect without the risk of iodine penetrating the corneal stromal layers and damaging corneal fibroblasts;

(iii) before considering the introduction of more powerful antibiotics in the storage solution, careful monitoring of contamination levels during the storage process will reveal whether reduction can be expected from other collection methods, improved working conditions, and repeated decontamination procedures.

The authors thank Mr B Willekens for performing the EDX investigations, the staff of the cornea bank for making tissue available and performing the culture experiments, the Department of Immunology for its kind gift of collagen I, the Department of Photography for preparation of the figures, and the Department of Microbiology, Academic Medical Centre, Amsterdam for determination of the contaminating microbes.


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