Evaluation of potential organ culture media for eye banking using human donor corneas

T Møller-Pedersen, U Hartmann, H J Møller, N Ehlers, K Engelmann

Abstract

**Aim**—To evaluate the ability of different commercially available cell culture solutions to preserve human donor corneas during 3 weeks of “closed system” organ culture at physiological temperature. This screening was performed in an attempt to establish a rational basis for the development of a serum-free organ culture medium for eye banking.

**Methods**—72 normal human donor corneas were organ cultured for 21 days at 31°C in eight different test media (nine corneas in each group). The basic culture solutions included: minimal essential medium (MEM), MEM with stabilised L-glutamine, M199, DIF-1000, SFM, F99, and F99 with ascorbic acid, insulin, bFGF, transferrin, selenium, and lipids (termed F99-Sr). All media were supplemented with 2% fetal calf serum (FCS), except for MEM, which was also studied at 8% FCS. The evaluation parameters included: (1) the endothelial cell loss as evaluated using trypan blue staining; (2) the ability of keratocytes and endothelial cells to incorporate tritiated uridine into RNA as evaluated using autoradiography and digital image analysis; (3) the leakage of immunogenic keratan sulphate as assessed using ELISA; and (4) changes in storage medium pH, glucose, and lactate content.

**Results**—SFM induced the lowest endothelial cell loss of 14% (SD 2%) and the highest RNA synthesis rates of all test media. Corneas stored in SFM also showed the least leakage of keratan sulphate as assessed using ELISA; and SFM appears to be the most promising medium (MEM), supplemented with varied FCS, except for MEM, which was also studied at 8% FCS. The evaluation parameters included: (1) the endothelial cell loss as evaluated using trypan blue staining; (2) the ability of keratocytes and endothelial cells to incorporate tritiated uridine into RNA as evaluated using autoradiography and digital image analysis; (3) the leakage of immunogenic keratan sulphate as assessed using ELISA; and (4) changes in storage medium pH, glucose, and lactate content.

**Conclusion**—Among the present test solutions, SFM appears to be the most promising candidate for a new corneal organ culture medium and should be further tested and possibly refined to effectively substitute serum addition.

More than 14 000 human donor corneas are processed annually in the European eye banks. Organ culture is the preferred storage method, in which the cornea can be kept at 31–37°C for a current maximum of 7 weeks and grafted successfully afterwards. During the organ culture period, the donor tissue undergoes screening for microbial contamination and positive serology (hepatitis, HIV, etc) as well as optional HLA typing and matching. The current preservation media generally consist of a basic culture solution, usually minimal essential medium (MEM), supplemented with varying amounts of fetal calf serum (FCS) ranging from 2% to 10%. Currently, this FCS enrichment is mandatory because of a lack of many essential (and unidentified) components in the basic culture media. Unfortunately, there is an unpredictable variation in the ingredients and composition of different FCS batches, which appears to be closely related to variation in the postoperative graft quality and performance. Furthermore, there is a potential risk of transferrable diseases and other unwanted side effects from unknown hazardous components in bovine serum. It is therefore desirable for the European eye banks to search for more standardised and optimised organ culture conditions using well characterised and serum free preservation media.

For these reasons, we have initiated experimental studies to evaluate systematically the impact of a wide range of commercially available basic culture solutions (which may work as starting points for subsequent refinements). Based on the ability to induce proliferation and morphological changes in primary cultures of human corneal endothelial cells (HCEC), we have recently performed an initial evaluation and comparison of 11 different cell culture media supplemented with varying concentrations of FCS ranging from 0% to 10%. Since this HCEC growth assay may not be a fully relevant screening model for the entire cornea, it is important to perform additional analyses of the storage media quality using intact donor tissue. Therefore in the present study, a total of 72 normal human donor corneas were obtained post mortem and organ cultured for 21 days in eight different test solutions. This paper summarises our evaluation of these media using quantitative viability parameters.

**Materials and methods**

**Organ culture technique**

A total of 72 human donor eyes (all without known corneal disease) were obtained post mortem from the cornea banks of Århus, Denmark and Hamburg, Germany. Mean donor
Figure 1 Autoradiograph of human donor endothelial cells following 48 hours of 3H-uridine incorporation during corneal organ culture. (A) Dense accumulations of silver grains cover three radiolabelled nuclei undergoing RNA synthesis (haematoxylin and eosin counterstain). Profiles of the same nuclei (B) and the area of the autoradiographic silver grains covering these profiles (C) as assessed by digital image analysis.

<table>
<thead>
<tr>
<th>Name</th>
<th>Basic component</th>
<th>% Serum supplementation</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>MEM-8%</td>
<td>MEM-Earle (Seromed)</td>
<td>8</td>
<td>Commonly used for organ culture in European eye banks^1</td>
</tr>
<tr>
<td>MEM-2%</td>
<td>MEM-Earle (Seromed)</td>
<td>2</td>
<td>Commonly used for organ culture in European eye banks^1</td>
</tr>
<tr>
<td>MEM-G</td>
<td>MEM-Earle (Seromed)</td>
<td>2</td>
<td>MEM supplemented with 2 mM l-analyl-l-glutamine, a stabilised form (dipeptide) of the amino acid l-glutamine</td>
</tr>
<tr>
<td>F99</td>
<td>M199 + Ham's F12 (1:1) (Gibco)</td>
<td>2</td>
<td>Used for corneal endothelial cell growth^11</td>
</tr>
<tr>
<td>F99-Sr</td>
<td>M199 + Ham's F12 (1:1) (Gibco)</td>
<td>2</td>
<td>F99 supplemented with ascorbic acid, insulin, bFGF, transferrin, selenium, and lipids^14</td>
</tr>
<tr>
<td>M199</td>
<td>M199-Earle (Seromed)</td>
<td>2</td>
<td>Intended for short term corneal preservation at 4°C (cold storage for up to 10 days)</td>
</tr>
<tr>
<td>SFM</td>
<td>Company formula (Gibco)</td>
<td>2</td>
<td>Developed for serum free growth of vascular endothelial cells</td>
</tr>
<tr>
<td>DIF-1000</td>
<td>Company formula (Seromed)</td>
<td>2</td>
<td>Initially used for serum free growth of hybridoma cells</td>
</tr>
</tbody>
</table>

See company information sheets for further details on the specific composition of each medium.

Age was 68 (SD 13) years (range 24–89) and the mean postmortem time was 39 (13) hours (range 23–72) (defined as the interval between death and incubation of the cornea). All corneas included in this study fulfilled the standards of the European Eye Bank Association for donor material used for grafting. The eyes were rinsed under tap water and the corneas were carefully excised including a 2 mm scleral rim. The corneas were randomly divided into eight experimental groups (with nine corneas each) and incubated at 31°C for 21 days (hanging in a suture through the scleral rim) in 70 ml airtight glass bottles containing 20 ml atmospheric air and 50 ml of one of the eight test media (Table 1). These media were selected following an initial evaluation using primary cultures of HCEC as recently reported. Different supplements were added to ensure that all storage solutions contained: 12.5 mM HEPES, 2.0 mg/ml gentamicin, and (in seven of the eight media) 2 mM of the amino acid l-glutamine that is known to be unstable in solution. Therefore, one group of corneas (termed MEM-G) received 2 mM l-analyl-l-glutamine (Table 1), which is a stabilised form (dipeptide) of l-glutamine. Furthermore, 2% FCS (batch B-425C, Seromed) was added to seven of the eight test groups, whereas one group of corneas (termed MEM-8%) received 8% FCS supplementation (Table 1). Finally, pH was adjusted to 7.4 in all culture media using NaHCO₃. To maintain sterility, the media were not changed during the 21 day “closed system” organ culture.

ASSESSMENT OF ENDOTHELIAL CELL DENSITY
Immediately before and after incubation, the endothelial cell density of each donor cornea was estimated using trypan blue staining and osmotically induced dilatation of the intercellular spaces by 0.9% NaCl. The difference between the two counts represents the cell loss during the storage period.

ESTIMATION OF RNA SYNTHESIS
To radiolabel the RNA synthesis, 1.0 µCi/ml 5,6-3H-uridine (specific activity 38.5 Ci/mmol, NEN Research Products) was added to each bottle 48 hours before termination of the incubation as described previously. At day 21, central pieces of the corneal endothelium including Descemet’s membrane were gently peeled off, mounted on gelatinised slides, fixed in formalin, air dried, and prepared for autoradiography. Similarly, samples of the central stroma were fixed in formalin and processed for histology and autoradiography. Using the same batch of reagents, all slides were assayed in parallel, including appropriate control sections from corneas incubated without radioactivity.

Following photographic development (at 4°C for 11 days) and a haematoxylin and eosin counterstain, 30 nuclear profiles of keratocytes and endothelial cells, respectively, were randomly selected and digitised for each cornea (Fig 1A). In all images, the area of the nuclear profiles (Fig 1B) and the area of autoradiographic silver grains covering the profiles (Fig 1C) were calculated using the same automated algorithms from the software package Matrox Inspector Version 1.5. For each cornea, the percentage of silver grains on the nuclear area represents a quantitative measure of the RNA synthesis rate that can be used as a simple estimate of cell viability. Obviously, the dose-response relation for this assay may not be fully linear because of saturation of the photographic emulsion.

IMMUNOGENIC KERATAN SULPHATE
Before and after incubation, all storage media were assayed for their content of immunogenic keratan sulphate (KS) using the monoclonal antibody 5D4 (from ICN, code no 69-625-2) in a sensitive, competitive ELISA.10 11 This
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5D4 antibody recognises an epitope on the sulphated polysaccharide structure of KS proteoglycan and has previously been used to detect KS in the cornea and other tissues. The difference between the two measurements represents the leakage of stromal KS during the storage period and is expressed in µg/ml medium.

METABOLIC PARAMETERS

Before incubation and at day 21, duplicate samples of all culture media were obtained for measurements of glucose, lactate, and pH as previously reported.

STATISTICS

Data are given as mean (SD). Using SIGMASTAT Version 2.0 (Jandel Corporation, CA, USA), differences in the mean values among the groups were compared by one way analysis of variance and Student-Newman-Keuls method for all pairwise comparisons (within each column) revealed the following significant differences (p<0.05): (a versus b, c, d, e, f); (f versus all); (g, k, l versus all); (h versus i, j); (m, n, o versus p, r, t); (q versus t) (s versus p, q, r, t); (y versus all except u, x); (u versus v); (x versus v, z).

RESULTS

LOSS OF ENDOTHELIAL CELLS

Before incubation, the average endothelial cell density for all 72 corneas was 2801 (327) cell/mm² (range 2210–3680) with no significant differences between the groups. Following 21 days, the endothelial cell loss averaged 17.5% with considerable variation among the media. Corneas stored in MEM-8% showed the lowest cell loss, 11% (4%), compared with corneas kept in MEM-2%, 19% (6%) (p<0.05), demonstrating a marked protective effect of a higher serum level (Table 2). Among media with 2% FCS supplementation, SFM showed the lowest cell loss, 14% (2%), whereas the highest cell loss, 29% (4%), was observed in DIF-1000.

KERATOCYTE AND ENDOTHELIAL CELL RNA SYNTHESIS

After 21 days, a significant correlation (r = 0.70, p <0.05) was found between the RNA synthesis rates (degree of autodiagnostic silver grain staining) of keratocytes and endothelial cells; indicating that the two cell types have parallel protein synthesising activities during storage. However, profound variation was observed among the test media. The highest level of silver grain staining (38–57%) was found in MEM-8% and in SFM, followed by MEM-2% and MEM-G (29–37%) (Table 2). An intermediate silver grain staining of 19–29% was seen in F99, F99-Sr, and M199, whereas DIF-1000 revealed the lowest RNA synthetic activity (4–15% silver grains) with many unlabelled cells (Table 2). For both keratocytes and endothelial cells, the degree of staining was found to be inversely correlated with the loss of endothelial cells, r = −0.63, p <0.05 and r = −0.73, p<0.05, respectively. In other words, the lower the capacity to synthesise RNA (and thereby to renew critical proteins), the higher the level of endothelial cell death.

LEAKAGE OF KERATAN SULPHATE

Before incubation, the content of KS was below the detection limit in all test media. After 21 days, the KS concentration in the storage media varied from 22 to 290 µg/ml with substantial variation among the groups. The lowest KS loss of 57 (24) µg/ml was detected in SFM, followed by 77 (28) µg/ml in MEM 8%, and 77 (28) µg/ml in M199 (Table 2). For the remaining five media, the average KS loss ranged from 118 to 144 µg/ml, which corresponds to approximately 1.5% of the total content of immunogenic KS in human donor corneas. Interestingly, the storage medium content of KS correlated inversely with both the endothelial cell density at day 21 (r = −0.45, p<0.05) and with the endothelial cell RNA synthesis rate (r = −0.49, p <0.05); suggesting that stromal KS leakage may depend directly on the integrity and barrier function of the endothelium as previously observed.

METABOLIC CHANGES IN THE TEST MEDIA

In all test solutions, the initial metabolic characteristics resembled that of a physiological system with pH = 7.4, a lactate concentration ranging from 0.3 to 1.3 mM, and (in six of the eight media) a glucose concentration ranging from 5.5 to 8.0 mM. However, two media (SFM and DIF-1000) had a high initial glucose content of about 17 mM. Interestingly, corneas stored in these two media displayed a significantly (p<0.05) higher glucose utilisation (~6.9 to ~7.3 mM) during the 21 day incubation compared with all other test solutions (~4.3 to ~5.5 mM) (Table 3). This finding suggests that a general lack of glucose may limit the metabolic capacity of donor corneas kept in MEM, M199, and F99 based media.

Table 2 Characteristics of human donor corneas following 21 days of organ culture in eight different storage media

<table>
<thead>
<tr>
<th>Test medium</th>
<th>Endothelial cell loss (%)</th>
<th>Keratocyte RNA synthesis (% silver grains per nuclear area)</th>
<th>Keratan sulphate loss (µg/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MEM-8%</td>
<td>11 (4)a</td>
<td>57 (12)g</td>
<td>74 (41)u</td>
</tr>
<tr>
<td>MEM-2%</td>
<td>19 (6)b</td>
<td>29 (11)</td>
<td>127 (63)</td>
</tr>
<tr>
<td>MEM-G</td>
<td>17 (4)c</td>
<td>37 (10)h</td>
<td>118 (63)</td>
</tr>
<tr>
<td>F99</td>
<td>16 (4)</td>
<td>21 (7)</td>
<td>144 (60)v</td>
</tr>
<tr>
<td>F99-Sr</td>
<td>17 (3)e</td>
<td>19 (11)</td>
<td>132 (34)</td>
</tr>
<tr>
<td>M199</td>
<td>14 (2)</td>
<td>48 (8)</td>
<td>57 (24)r</td>
</tr>
<tr>
<td>SFM</td>
<td>29 (4)</td>
<td>15 (5)</td>
<td>139 (40)x</td>
</tr>
</tbody>
</table>

Data are mean (SD), n=9.

One way analysis of variance and Student-Newman-Keuls method for all pairwise comparisons (within each column) revealed the following significant differences (p<0.05): (a versus b, c, d, e, f); (f versus all); (g, k, l versus all); (h versus i, j); (m, n, o versus p, r, t); (q versus t) (s versus p, q, r, t); (y versus all except u, x); (u versus v); (x versus v, z).

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should be noted that SFM showed the highest lactate production of 12.4 mM (p < 0.05), whereas corneas kept in DIF-1000 revealed the lowest level of lactate formation, 6.9 mM (Table 3). Assuming a direct equivalence of 1 M glucose to 2 M lactate (and ignoring other metabolic pathways), it appears that corneas kept in SFM predominantly utilises glucose by 85% glycolysis and 15% oxidation (Table 3), which mimics the metabolism of a normal cornea. Comparable glycolytic activity (72–88%) was observed in all other test media, except for DIF-1000 that only showed 54% glycolysis (p < 0.05) (Table 3). It is interesting to note that the accumulation of lactate was inversely correlated with the endothelial cell loss (r = −0.43, p < 0.01), confirming that a well functioning cornea has a high glycolytic activity. As for changes in storage medium pH, the eight groups showed a significant variation ranging from −0.37 to −0.67 units (Table 3); however, no systematic impact of these pH changes could be identified on neither the endothelial cell loss, the RNA synthesis rates, nor the leakage of keratan sulphate.

DONOR CHARACTERISTICS

No systematic influence of neither donor age, sex, medication, cause of death, nor the post mortem time could be identified on any of the study parameters.

Discussion

The present study showed no major differences between corneas stored in MEM and F99 based media, although F99 generally contains higher levels of multiple essential nutrients and supplements. Neither did the specific enrichment of F99 with ascorbic acid, insulin, bFGF, transferrin, selenium, and lipids show any improvement on corneal cell density, viability, or metabolism (Tables 2 and 3; compare MEM-G with MEM-2%). Thus, the endothelial cell loss occurred independently of the presence of labile L-glutamine in the storage media, which is consistent with a previous study of porcine corneas. Substitution of the labile amino acid L-glutamine with the stable dipeptide L-analyl-L-glutamine had no significant impact on the loss of endothelial cells, the RNA synthesis rates, or the leakage of keratan sulphate (Table 2; compare MEM-G with MEM-2%). Thus, the endothelial cell loss occurred independently of the presence of labile L-glutamine in the storage media, which is consistent with a previous study of porcine corneas.

The most viable and best preserved donor corneas generally showed a high glucose turnover with a glycolytic activity close to 85% (Table 3); which according to Riley mimics the metabolism of a normal cornea. In other words, a high lactate concentration in the storage medium appeared to be a simple marker of cell viability, that in many cases indicated a well functioning donor cornea with a low endothelial cell loss. By contrast, the observed change in storage medium pH (−0.37 to −0.67 units)
appeared to be a poor indicator of decreased cell density and viability. Thus, a reduced extracellular pH did not appear to limit corneal cell viability, although it may increase the metabolic stress on the tissue. This finding is in agreement with a previous study of organ cultured cat corneas showing a normal intracellular pH despite an extracellular pH decline of 0.6 units.16

In conclusion, the majority of the tested storage media (MEM-2%, MEM-G, M199, F99, and F99-Sr) showed comparable potentials for corneal organ culture at 2% FCS supplementation. However, there was a clear tendency for the medium SFM to maintain a higher endothelial cell density and viability at 2% FCS. Thus, SFM should be further evaluated in the absence of serum and possibly refined and optimised. In such studies, it might be beneficial not to use “closed system” organ culture but instead change the medium—for example, every week (“open system”) in order to maintain corneal cell viability under long term, serum free storage. Further evaluation of SFM will be of interest.

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