Bullous keratopathy: a study of endothelial cell morphology in patients undergoing cataract surgery

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SUMMARY Preoperative and postoperative endothelial cell morphology was examined in three types of patients participating in a randomised, controlled trial of cataract surgery. The first sustained modest endothelial cell loss and served as controls, the second and third sustained high cell loss, but only the latter progressed to bullous keratopathy. The technique of examining endothelial cell morphology is described and its validity assessed. Measures of cell area, number of sides, side length, and variation in cell size and cell shape were utilised. No significant differences were found between the three groups by any of the preoperative morphological measures. Furthermore, postoperative values of variation in cell shape and cell size did not differ significantly between the groups. No evidence was found in our patients to support the hypothesis that eyes at particular risk of developing bullous keratopathy may be differentiated on the basis of their preoperative or early postoperative endothelial cell morphology.

Bullous keratopathy has become the commonest condition requiring penetrating keratoplasty.1 2 The increase in incidence of bullous keratopathy may have resulted from the increased volume of cataract surgery, new surgical techniques, or increased implantation of intraocular lenses of varying design. The exact contribution of each potential cause has not been determined, but a better understanding is desirable to identify vulnerable cases and risk factors. Much emphasis has been placed recently on the morphology of the corneal endothelium,3 and a suggestion has been made that eyes developing bullous keratopathy may have more preoperative polymorphism or pleomorphism. The purpose of this paper is to report on an examination of endothelial cell morphology pre- and postoperatively of eyes developing bullous keratopathy and compare that to findings in controls.

Material and methods

Patients studied were participating in a randomised controlled trial of cataract surgery which has been in progress for six years, for five of which there was no loss to follow-up except through death. Detailed accounts of the trial design and surgical techniques have already been reported.4 The majority of patients were treated by one experienced surgeon who contributed 274 eyes of 268 patients with a mean age of 72 years and a range of 55–89 years. All the patients in the present study on bullous keratopathy contributed only one eye in the study and had only one eye operated on at the time of this assessment. Patients were randomised into one of three treatment groups which were:

Group A. Intracapsular extraction followed by contact lens fitting.

Group B. Intracapsular extraction and implantation of an iris supported lens—Federov I.

Group C. Extracapsular extraction and implantation of a Binkhorst two-loop iridocapsular lens.

Patients had endothelial cell photographs taken preoperatively, at one month, six months, and yearly after surgery. Cell densities were derived in all patients by a counting method which masks the patients' identity. The method of photography5 and precision estimates6 have been reported.

A detailed protocol and structured follow-up enabled us to identify three groups of patients. The first group were patients who underwent uneventful surgery and postoperative recovery and who sustained only modest endothelial cell loss (mean cell loss six months after operation=10.03%, SD=6.56%). This group served as controls. The second group were patients whose surgery and/or
postoperative recovery was complicated and who sustained large cell loss in the early postoperative period but did not develop bullous keratopathy (mean cell loss six months after operation=33.66%, SD=18.2%). The third group sustained large endothelial cell loss and developed bullous keratopathy (mean cell loss six months after operation=31.68%, SD=16.9%).

Preoperative and one month postoperative endothelial cell photographs from these three groups of patients were enlarged to allow analysis of cell morphology. The original magnification factor of 148 times was increased to 311 times and photographic contrast enhancement techniques were used in developing the film. The apices of the endothelial cells were then digitised on a Summagraphics MM 1201 digitising pad with a digitising pen and the data analysed by an IBM personal computer. Digitisation in this context means the characterisation of points by a set of numbers. The digitising tablet employed was able to determine the location of points ±0.625 mm with repeatability for the same point of ±0.25 mm or better (manufacturers’ specification). For most photographs 75 cells were digitised and in others a minimum of 50 cells. Evidence for the adequacy of a 50-cell sample has been presented by Bourne.7

A program was developed which, from digitisation of the apices of endothelial cells, will compute the area of individual cells, that of the smallest and largest cells, a range of cell areas, side lengths and the number of sides, and the perimeter of each cell. From these basic data two further measures were computed. As an index of variation in cell area the coefficient of variation was calculated by dividing the standard deviation of the cell area by the mean cell area and then expressing this as a percentage. This ratio is independent of cell size and provides a quantitative measurement of variation in cell area (polymegathism). Variation in cell shape was quantified by comparing the digitised cells with equilateral hexagons. For an equilateral hexagon the ratio of area:perimeter is a constant (sin 60°/12 or 0.07217). The deviation of digitised cells from hexagonality may be expressed as a percentage according to the equation [(A/p^2)/0.07217]×100, such that, if the cell is an equilateral hexagon, the index will be 100. This value is a dimensionless expression of variation in cell shape (pleomorphism) which is independent of size and orientation.8

**Table 1 Validation study**

<table>
<thead>
<tr>
<th>Digitisation procedure</th>
<th>Mean cell area of large cells</th>
<th>Mean cell area of small cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>50 Cells from 2 photos five times</td>
<td>287.85±2.9</td>
<td>134.10±1.8</td>
</tr>
<tr>
<td>3 Central photos of 2 patients</td>
<td>SD=1.0% of mean</td>
<td>SD=1.3% of mean</td>
</tr>
<tr>
<td></td>
<td>292.93±3.4</td>
<td>119.33±1.9</td>
</tr>
<tr>
<td></td>
<td>SD=1.2% of mean</td>
<td>SD=1.6% of mean</td>
</tr>
</tbody>
</table>

five times, one with a population of large cells and the other small cells. Thus five cell areas were obtained for each photograph.

Reproducibility was examined by digitising three specular microphotographs of the central corneal endothelium from two patients, again one with large and the other with small endothelial cells.

**Results**

The validation study showed a high degree of accuracy, precision, and reproducibility. Accuracy: the mean value of digitising 50 hexagons of calculated area five times differed from the calculated area by only 3.15%. Precision: repeated digitisation of 50 cells from a sample of large and a sample of small cells produced a standard deviation of 1.0 and 1.3% of the mean (Table 1). As expected, the value was smaller for large cells. The mean cell area could be measured with a precision of 1.15% by this technique. Reproducibility: the variation of mean cell area between the three photographs of the same patient was about 1.2–1.6% (Table 1).

The results of the morphological analysis of preoperative photographs from the three groups of eyes are shown in Table 2. There was no significant difference in the values of the different features except for mean cell area. The mean area of the smallest cells in eyes developing bullous keratopathy was significantly larger, probably through inclusion of two eyes with low cell density.

One month postoperatively, as expected, the values of all the features examined were significantly greater in the two groups, with high cell loss in comparison with the control group. However, neither coefficient of variation nor deviation from hexagonality showed any significant change. In a few patients the photographs were not of good enough quality for digitisation.

**Discussion**

The validation study showed a high degree of accuracy, precision, and reproducibility. Our results on precision and reproducibility compare very favourably with those reported by others.7-11
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Table 2  Morphological analysis of preoperative photographs

<table>
<thead>
<tr>
<th></th>
<th>Controls n=15</th>
<th>High cell loss, patients n=7</th>
<th>Bullous keratopathy, patients n=9</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean cell area</td>
<td>427.66 M±5.56</td>
<td>430.94 SD=5.66</td>
<td>494.75 SD=10.3</td>
</tr>
<tr>
<td>Mean cell density</td>
<td>2802</td>
<td>2534</td>
<td>2483</td>
</tr>
<tr>
<td>Mean area of smallest cell</td>
<td>180.71 SD=7.81</td>
<td>170.81 SD=23.59</td>
<td>233.82 SD=61.51</td>
</tr>
<tr>
<td>Mean area of largest cell</td>
<td>855.08 SD=6.56</td>
<td>865.76 SD=93.59</td>
<td>933.59</td>
</tr>
<tr>
<td>Mean no. of sides per cell</td>
<td>6.01 SD=0.14</td>
<td>5.87 SD=0.14</td>
<td>6.00</td>
</tr>
<tr>
<td>Mean side length</td>
<td>13.0±7</td>
<td>13.5±3</td>
<td>14.16</td>
</tr>
<tr>
<td>Coefficient of variation</td>
<td>31.09 SD=0.96</td>
<td>30.41 SD=28.58</td>
<td>28.58</td>
</tr>
<tr>
<td>Deviation from hexagonality</td>
<td>92.98 SD=1.43</td>
<td>92.26 SD=93.51</td>
<td>93.51</td>
</tr>
</tbody>
</table>

Cell density ranges 1333-1186 1260

*Areas in μm²; density in cells/mm²; length in μm.
*p<0.05 vs controls. †No significant difference from controls.

Preoperative assessment of endothelial cell morphology showed values for mean cell area consistent with typical endothelial cell densities for this age group of patients (for example, for controls= 2338 cells/mm²).¹² Hexagons were the most commonly observed cell shape as shown by the mean number of sides—close to six observed in each group (Fig. 1). The values for the coefficient of variation were within the range previously quoted for patients obtained before cataract surgery¹ and suggest that cells were relatively uniform in size. Similarly values for deviation from hexagonality were all very high, suggesting homogeneity of cell shape.

Interestingly, there was no significant difference between the three groups on any of the preoperative morphological measures studied except mean area of the smallest cell area between controls and bullous keratopathy (BK) patients. This is due to the fact that two of the eyes with BK had a preoperative density of <2000 cells/mm². It has been suggested that cellular polymegathism and pleomorphism reflect susceptibility to surgical trauma¹³ and in particular postoperative corneal oedema.¹ However, our results suggest that for eyes without pre-existing disease preoperative endothelial morphology does not differentiate either patients who will undergo large endothelial cell losses or those who will progress to bullous keratopathy from controls.

Postoperative analysis of endothelial cell morphology revealed expected increases in mean cell area as well as area of the smallest and the largest cells in all groups, with the appearance of cells with the greater number of sides in the high cell loss groups (Table 3 and Fig. 2). However, while cells increased in size, the fairly constant values for coefficient of variation and deviation from hexagonality suggest that in general they increased uniformly, so that variation in size and shape between cells changed little. It is just possible that at one month after surgery central cellular rearrange-

![Fig. 1  Frequency distribution of the number of sides of the cells in the three groups of eyes before operation.](image)

Table 3  Morphological analysis of postoperative photographs

<table>
<thead>
<tr>
<th></th>
<th>Controls n=12</th>
<th>High cell loss, patients n=6</th>
<th>Bullous keratopathy, patients n=5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean cell area</td>
<td>498±20</td>
<td>923.03±32</td>
<td>885.83±52</td>
</tr>
<tr>
<td>Mean cell density</td>
<td>2436</td>
<td>1294</td>
<td>1602</td>
</tr>
<tr>
<td>Mean area of smallest cell</td>
<td>244±6 SD=269</td>
<td>212 SD=423</td>
<td>432.34±32</td>
</tr>
<tr>
<td>Mean area of largest cell</td>
<td>35±6 SD=120.01</td>
<td>510 SD=34</td>
<td>186.03</td>
</tr>
<tr>
<td>Mean no. of sides per cell</td>
<td>9.57±9 SD=0.10</td>
<td>6.21±9</td>
<td>6.07</td>
</tr>
<tr>
<td>Mean side length</td>
<td>14±31</td>
<td>18.61±3</td>
<td>18.86±3</td>
</tr>
<tr>
<td>Coefficient of variation</td>
<td>28.55 SD=3.58</td>
<td>5.01 SD=2.56</td>
<td>26.28±9</td>
</tr>
<tr>
<td>Deviation from hexagonality</td>
<td>92.37 SD=1.21</td>
<td>92.55±9</td>
<td>92.19±9</td>
</tr>
</tbody>
</table>

Cell density ranges 707-551 133

*Areas in μm²; density in cells/mm²; length in μm.
***p<0.001
**p<0.01
*p<0.05 vs controls. †No significant difference from controls.
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References


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