Corneal endothelial cell abnormalities in an early stage of the iridocorneal endothelial syndrome

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Abstract
A corneal disc, obtained from a 52-year-old woman suffering from an early stage of the iridocorneal endothelial syndrome (ICE), was investigated by various morphological techniques to analyse the structural variations in the endothelial cells and to identify the collagen types within the abnormal layer of Descemet's membrane. Scanning electron microscopy of the posterior corneal surface revealed a mosaic of (a) flat hexagonal cells resembling irregular but normal endothelial cells, and (b) rounded hexagonal (ICE) cells with numerous surface microvilli. Degenerative changes were present in each cell type, but were more common in the flat hexagonal cells which contained intracytoplasmic spaces. By transmission electron microscopy the flat hexagonal cells exhibited many of the features of normal endothelial cells in terms of organelles and intercellular attachments, but lateral invaginations were absent. The ICE cells differed in that the apical surface was covered by microvilli and the cytoplasm contained tonofilaments, which were also observed by light microscopic immunocytochemical staining. Most commonly, intercellular attachments were rudimentary in both types of cell and intercellular spaces were dilated, but desmosomes were sometimes prominent in the ICE cells where interdigitations were pronounced. In some sectors, the basal surface of the ICE cells was indented by deposition of clumps of fibrillar collagenous material. An immunocytochemical study of the abnormal posterior deposits localised type IV collagen to the amorphous matrix and collagen types III and V, but not type I, to the collagen fibril bundles. Mononuclear inflammatory cells were identified between the ICE cells in the monolayer. The evidence suggests that some of the flat hexagonal cells were undergoing a degenerative change while others were transforming into ICE cells.

By transmission electron microscopy ICE cells have been shown to have endothelial cell characteristics. By contrast, ICE cells have been shown to have epithelial characteristics with desmosomes and intracytoplasmic tonofilaments. Descemet's membrane which normally contains two collagen types (IV and VIII) is thickened in ICE by the addition of a broad posterior collagenous layer (PCL) occasionally containing wide banded collagen. Small foci of fibrillar deposits behind the posterior non-banded zone were reported in one example of the early stage of the disease. An attempt to identify the collagen types in the PCL has not to our knowledge been attempted. In the present morphological study we describe the features of the endothelium in an early uncomplicated case of the ICE syndrome in which, with the application of immunogold immunocytochemistry, we have identified some of the constituents of the abnormal posterior collagenous layer which is secreted by the ICE cells.

Clinical history
A female patient (born 10 May 1941) attended
the Tennent Institute in January 1993 with a unilateral decrease in visual acuity which was associated with some diurnal fluctuation in vision, the vision being worst in the morning and clearing slightly by mid-day. Such symptoms had persisted for 5 years. There was no family history of corneal disease. The referring ophthalmologist considered that the anterior chamber was shallow and that a few peripheral anterior synechiae were present. On examination the vision in the affected eye was reduced to 6/24 with correction. There was stromal and epithelial corneal oedema and a distorted eccentric pupil with ectropion uveae. Fine endothelial detail was obscured by the oedema. No other features were distinguishable and in particular the other eye was entirely normal. Intraocular pressure in both eyes was within normal limits. Endothelial specular photomicroscopy (ESP) was performed with considerable difficulty because of the corneal oedema. In some areas, cells with central highlights were visible. Clinically this was felt to be ICE syndrome because of the unilaterality, the ESP appearance and the lack of any clinical features suggestive of posterior polymorphous dystrophy – namely, vesicles, snail track, and thickened Descemet’s membrane. The small ectropion uveae was also felt to be consistent with the ICE syndrome.

A corneal graft was performed on 12 January 1993 and at follow up 9 months later the graft has remained clear and the intraocular pressure has remained normal.

Materials and methods

The keratoplasty disc measured 8 mm in diameter and the stroma was cloudy. The specimen was divided into four, and three parts were fixed in glutaraldehyde (2%) for paraffin histology, scanning electron microscopy, and conventional transmission electron microscopy. The remaining part was fixed in a paraformaldehyde (4%)glutaraldehyde (0.5%) mixture for 2 hours at room temperature for immunogold immunocytochemistry.

Paraffin sections were stained with haematoxylin and eosin, periodic acid Schiff, and Masson and were also used for an immunohistochemical study of cytokeratins in the epithelium and the endothelium. The following antibodies were applied – CAM 5-2, AE 1/3, NCL, and MNF. The tissue for transmission electron microscopy was processed through to Araldite and semithin sections were stained with toluidine blue. Ultrathin sections were examined in a Jeol 1200 EXII transmission electron microscope. The quadrant used for scanning electron microscopy was subject to critical point drying before gold coating and examination in a Jeol JSM 64000 scanning electron microscope. The block allocated for immunocytochemistry was processed for London resin (LR) white embedding. Details of the techniques for embedding, antibody labelling, and immunocytochemistry have been provided elsewhere.

Results

LIGHT MICROSCOPY

The epithelium was oedematous and the basement membrane was multilayered and infolded in parts: there was no evidence of a band keratopathy. Bowman’s layer was intact and there were plentiful stromal keratocytes, but in one half of the cornea the lamellae appeared to be more widely dispersed owing to oedema. Descemet’s membrane measured 6 µm but there was evidence of multilayering in the PAS stained sections (Fig 1). The normal cuboidal endothelium was replaced either by vacuolated cells or rounded cells and some cells were remarkably attenuated (Figs 1a, 1b). Mononuclear inflammatory cells were present within the monolayer (Fig 1b). Consistent results were obtained with the immunohistochemical markers for cytokeratins. The rounded ICE cells stained positively while the flattened and vacuolated cells were weakly staining (Fig 1d). The epithelium stained positively for each of the antibodies used.

Figure 1. Light micrographs of the posterior corneal surface. (a) Vacuoles (arrowheads) were a prominent feature in some areas of the endothelium. (b) In other sectors the cells were rounded and inflammatory cells (arrowheads) could be identified in the monolayer. (c) In the periodic acid Schiff stained paraffin section, Descemet’s membrane (arrowheads) appears bilayered and the endothelial cells are vacuolated or attenuated. (a) and (b) Araldite toluidene blue, ×630; (c) paraffin ×750. (d) Some of the cells (corresponding to rounded cells) on the posterior surface stain positively with cytokeratin markers while other cells (corresponding to the flat vacuolated cells) are negative (CAM 5-2).
Figure 2 Features of the posterior corneal surface as seen by scanning electron microscopy. (a) Low power view of the flat hexagonal cells: note the degenerating cells (arrowheads). (b) Higher power to contrast the surface of hexagonal cells with that of the ICE cells which are covered by microvilli: note cilia on the surface of the hexagonal cells (arrowheads). (c) The ICE cells sometimes showed widening of the intercellular spaces (arrow): note ruptured cyst in the hexagonal cells (arrowhead). (d) The ICE cells have prominent nuclear bulges and cilia (arrowheads). ((a) x800; (b) x5000; (c) x1600; (d) x1400.)

SCANNING ELECTRON MICROSCOPY
The appearance of the cells on the posterior corneal surface varied considerably. Small areas of flat cells resembled the normal hexagonal endothelium, but there was marked variation in shape and diameter (from 10 to 20 μm) of these cells (Fig 2). Cilia were present and the edges of the cells were lined by microvilli (Fig 2b). Ruptured cystic blebs exposed the cytoplasmic contents and extruded cytoplasmic debris lay on the surface of the adjacent hexagonal cells.

The majority of the cells were more regular in shape and size being hemispherical and covered by numerous microvilli: these cells are conventionally referred to as ICE cells. The microvilli were fewer in the central part of the cell, where there was a bulge and here a central cilium was easily identified. In other areas intercellular spaces were pronounced. Considerably fewer ruptured cysts were observed within hemispherical cell populations than among hexagonal cells. Binucleate cells were not seen.

TRANSMISSION ELECTRON MICROSCOPY

Hexagonal cells
These were identified as endothelial cells by the smooth posterior surface, typical mitochondria with tubular cristae, cilia, and a paucity of cytoplasmic filaments. Some cells contained isolated melanosomes (Fig 3) and areas of intracytoplasmic rarefaction and, in some groups of cells, what appeared to be large intracytoplasmic cysts were found. Intercellular lateral infoldings were not pronounced, but the junctions were those of the normal endothelium: occasionally the intercellular space was widened (Fig 3).
Hemispherical (ICE) cells
These were characterised by numerous microvilli on the posterior surface, abundant intracytoplasmic filaments (80–100 nm diameter) (Fig 4) and mitochondria with lamellar cristae (Fig 5). Intercellular indentations were sometimes prominent and desmosomal attachments were an occasional finding. Some of the cells contained an isolated melanosome. Inflammatory cells (mainly lymphocytes and a few macrophages) were present within the monolayer. Descemet’s membrane was thickened beneath the ICE cells by irregular deposition of clumps of striated fibrils embedded in an amorphous matrix: these nodules also projected into the cells (Fig 6). Wide banded collagen was absent.

Figure 4 Degenerative changes in the ICE cells were seen as enlarged spaces within the cytoplasm and between the cells (arrowhead, shown in (a)) and numerous pinocytotic vesicles adjacent to the microvilli (arrow, shown in (b)). Note the prominent intracytoplasmic filaments in the cells. (Aurilidse (a) ×2000; (b) ×10 000.)

IMMUNOGOLD IMMUNOCYTOCHEMISTRY
Collagen types I, III, and V were localised to striated collagen fibrils in Bowman’s layer and in the corneal stroma (Fig 7). Labelling for type IV collagen was present in the posterior non-banded zone of Descemet’s membrane (Fig 8), but was absent from the basement membrane of the corneal epithelium. No labelling for type II collagen was seen in any of the regions studied. The localisation of collagen types I–V in the present specimen, excepting Descemet’s membrane (see below), was therefore identical to that seen in immunogold studies of aged human cornea.14–16

The accumulations of striated collagen fibrils which were observed between the ICE cells and Descemet’s membrane were arranged either in whorls which invaginated the adluminal surface of the cells (Figs 8, 9) or in long thin bundles lying parallel to the endothelial surface (Fig 6). Both arrangements of fibrils exhibited intense labelling for collagen types III and V (Fig 9) and type IV was present in the enveloping amorphous matrix (Fig 8). Labelling for types I, II, and IV collagen was consistently absent from such fibril accumulations.

Immunogold particles were rarely observed in the rabbit antimouse immunogold control sections (omission of primary antibody) and were sparse in the normal goat serum control sections (that is, substitution of the primary antibody with non-immune serum from the same species in which the primary antibody was raised).

Discussion
Many of the cases reported as the ICE syndrome have been taken from patients suffering from abnormalities additional to those in the cornea. The value of the present case lies in the fact that it represents the disease at an early and uncomplicated stage.

In the present case, light microscopy was of value in confirmation of the clinical diagnosis of the ICE syndrome, because the flat vacuolated endothelial cells were easily distinguished from the humped ICE cells and immunocytochemistry identified cytokeratins in the latter cells. The focal positivity and minor variations we observed were also noted by Kramer et al.,16 while more extensive staining was reported by Hirst et al.16:
these differences may reflect the change from flat hexagonal cells to ICE cells as the disease progresses and at the end stage multilayering has been observed within the population. The presence of vacuolation in a distinct monolayer over a relatively thin Descemet’s membrane is dissimilar to many of the cases described in the ICE literature, particularly in the reports of Chandler’s syndrome in which Descemet’s membrane was thickened and the cells were attenuated and/or duplicated. Another discrepancy is that cystic blegging (Fig 2) was more common in the hexagonal cells compared with the ICE cells in the present case, rather than in the ICE cells as has been reported by others. A thickness of 6 μm of Descemet’s membrane was a surprising feature in the present case since in all of the other published reports, Descemet’s membrane has been markedly thickened. By light microscopy it was also possible to identify mononuclear inflammatory cells within the endothelium, but this feature, as these authors have stated, can only be regarded as supplemental to the diagnosis and is regarded as non-specific in other corneal disorders.

The ultrastructural appearances of the ICE cells within the endothelium. (a) A low power view to show cells with prominent microvilli. (b) A mononuclear inflammatory cell is present within the monolayer (arrows). (c) In some areas desmosomal attachments (arrowheads) are present on the interdigitating cell membranes. (d) Dense contractile filaments are present within the cytoplasm of some ICE cells. (a) × 2000; (b) × 10,000; (c) × 25,000; (d) × 100,000.)

Scanning electron microscopy of the abnormal endothelium in the ICE syndrome provides a great deal of additional information concerning variations in cell morphology. The differences in appearance of the flat hexagonal cells and the ICE cells in the present case were very similar to those previously reported in various disorders in the ICE group, with some cells exhibiting transitional features between the flat and microvillous surfaces. Not all cases classified as ICE in the literature have demonstrated a continuous monolayer and illustrations of isolated cells with extended processes on an exposed Descemet’s membrane have appeared. This extreme stage of endothelial degeneration and atrophy appears to be associated with extension of the endothelium over the chamber angle onto the iris surface and therefore to massive depopulation of the corneal endothelium as in Chandler’s syndrome. Depopulation is preceded by cell migration and this change is probably expressed as filopodial extension as seen by scanning electron microscopy, but this abnormality was not observed in other cases or in the present case. This substantiates the impression that flat hexagonal cells are present in the endothelium at the earliest stage of the disease.

One of the striking features of the hemispherical ICE cells is that they cannot be classified as typical endothelial or epithelial cells on the basis of their ultrastructural features. While
microvilli have been described in several papers\textsuperscript{10,12} and cytoplasmic blebbing in the report by Alvarado et al., only four authors have emphasised the presence of intracytoplasmic tonofilaments\textsuperscript{4, 10, 12} and desmosomal attachments,\textsuperscript{12, 13} which characterise the cells as epithelial. Prominent tonofilaments and desmosomal attachments between multilayered cells favour a diagnosis of posterior polymorphic dystrophy (PPD), but for this unequivocal diagnosis an identifiable inheritance pattern and bilaterality are required.\textsuperscript{4} It is possible that there is an overlap between PPD and the ICE syndrome and there has been considerable interest and speculation in the metaplastic potential of the corneal endothelium, which may be based on its origin from the neural crest.\textsuperscript{14}

The appearances of the dual cell population in the present case resembled in part the endothelial-type cells described by Alvarado et al\textsuperscript{10} and the epithelial-type cells described by Kramer et al.\textsuperscript{15} The flat hexagonal cells did not have the typical features of endothelial cells in terms of intercellular boundaries, but contained the characteristic mitochondria and cilia. The epithelial-type cells exhibited intercellular invaginations, but were united by desmosomal attachments and lined internally by microvilli.

These differences are regarded by morphologists as of considerable importance in ascertaining the histogenetic origin of a cell population, but in the case of the corneal endothelium, derived from neural crest cells with a specific potential for metaplasia, the commonly accepted rules for an ectodermal or mesodermal origin do not apply. The mixture of cytoplasmic membrane characteristics observed in the cells on the posterior cornea suggests that there is a transformation from hexagonal cells to ICE cells and that the viability of the endothelial cells is lower than that of the ICE cells.

The presence of inter- and intracytoplasmic blebs in the flat hexagonal cells is strong evidence in favour of a functional failure to control water movement. It is tempting to suggest that the transformation to ICE cells with complex interdigitations is an attempt to restore the functional integrity of the posterior monolayer. This would be a stress response and it is reasonable to assume that synthesis of a new posterior collagenous layer is part of that response. The presence of microvilli would increase the surface area of the cell membrane, so that this may be an aberrant attempt to control water movement. The presence of leaking and non-leaking areas of cells may be relevant to the unpredictable nature of corneal oedema in the ICE syndrome. A decrease in water movement across the cornea in the ICE syndrome has been demonstrated by Bourne and...
In the present specimen lymphocytes were found without difficulty in the endothelium among both the ICE cells and the relatively normal hexagonal cells. Inflammatory cells within the endothelium are a striking abnormality and have been recorded in two publications in addition to that of Alvarado et al. Rodrigues et al. suggested that an endotheliitis may be linked to a low grade iritis or that these cells are "incidental" wandering cells. Certainly the presence of melanosomes within the hexagonal cells and the ICE cells suggests that there had been some breakdown of the pigmented cells of the iris.

The mononuclear inflammatory cells, lymphocytes, and macrophages may be responsible for the degenerative changes, vacuolation, and cell death described in the endothelial cell population in other reports* and in the present case. In the present case it was apparent that the striated collagen fibres deposited presumably by the ICE cells contained type III and V collagen and the amorphous matrix contained type IV. Type I collagen is not normally found in Descemet's membrane,1 and type V collagen was located in the interfacial matrix between the stromal collagen and the anterior banded zone.2 It can therefore be concluded that the cells in the ICE syndrome were synthetically active but that the activity was aberrant in some aspects.

Cell culture experiments underline the latent ability of corneal endothelial cells to synthesise...
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the number of collagen types that are normally not found in Descemet’s membrane. Bosin corneal endothelial cells synthesise large amounts of types I, III, and V collagen when grown in cell culture (predominantly type III), whereas they synthesise types I and IV collagen when cultured on their Descemet’s membrane. It would appear that expression of type I collagen synthesis has independent regulation from that of III and IV. Therefore it is not surprising that ICE endothelial cells synthesised collagen types III and V without expression of type I collagen.

It is certainly relevant to note that polymorphonuclear leucocyte cells can modulate the synthetic activity of corneal endothelial cells by producing a corneal endothelial modulation factor (CEMF) that specifically transforms normal type IV collagen synthesising endothelial cells to type I synthesising cells. Since the inflammatory cells observed in this study and other studies were probably lymphocytes or macrophages this raises the interesting possibility that mononuclear inflammatory cells can produce similar CEMFs which can stimulate the ICE cells to produce an abnormal matrix.