Immunohistochemical analysis of lattice corneal dystrophies types I and II

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Abstract

Corneal buttons from four patients with lattice corneal dystrophy (LD) type I, thought to be an isolated corneal amyloidosis, and from six patients with LD type II, part of systemic familial amyloidosis, Finnish type (FAD; Meretoja’s syndrome), were studied by immunohistochemistry to determine the differential distribution in the amyloid deposits of amyloid P component (AP), mutated gelsolin specific for FAD, and native gelsolin. In both types of LD, antibodies to AP labelled lattice lines and a discontinuous layer of amyloid deposits under Bowman’s layer. In LD type II, particularly, they also reacted with stress-like amyloid deposits between corneal lamellae, eser. While the anti-FAD antiserum strongly labelled all amyloid deposits in LD type II, it failed to react unequivocally with them in LD type I. Both in LD type I and in two control specimens representing granular dystrophy, the monoclonal antibody (Mab) GS-2C4 to gelsolin faintly labelled some deposits, while in LD type II it reacted non-homogeneously with most amyloid deposits. In all specimens, MAb GS-2C4 labelled corneal epithelial cells and occasional stromal keratocytes and endothelial cells. The results suggest that Meretoja’s syndrome, a systemic disease, can be diagnosed even retrospectively from corneal buttons subjected to histopathological study.

Lattice corneal dystrophies (LD) are hereditary diseases characterised by lattice-like deposition of amyloid within the corneal stroma. Three clinically and histopathologically distinct types of LD have been reported. Types I and II are inherited as an autosomal dominant and type III as an autosomal recessive trait. Only LD type II is known to be associated with systemic amyloidosis, namely familial amyloidosis, Finnish type (FAD), also called Meretoja’s syndrome. LD is usually the earliest clinical finding in FAD. Slowly progressive cranial and peripheral neuropathy, dry and itchy skin and eyes, dermatohyalisis, depressed eyebrows, protruding lips, and intermittent proteinuria develop. Although it is most common in south eastern Finland, families have been encountered elsewhere in Europe and the United States.

Recently the genetic defect underlying FAD has been unravelled. The amyloid fibrils correspond to a 7 kDa internal degradation product of human gelsolin, a widely dispersed, calcium dependent, regulatory cytoplasmic and plasma protein involved in actin severing and gel sol transformation. It spans position 173 to 243 in the native protein, with asparagine 187 substituted for aspartic acid. This results from a single guanine to adenine transition at position 654, the first nucleotide of codon 187, within a highly conserved, repetitive motif of the gelsolin gene. FAD cosegregates with this mutation, it is particularly severe in those homozgyously affected, and the mutation has been shared by all Finnish and American patients examined.

Aberrant degradation of mutated gelsolin probably causes the amyloid deposition, as such peptides have a tendency to form amyloid fibres in vitro.

This report analyses the presence of native and mutated gelsolin in LD types I and II with Congo red and amyloid P component staining to determine whether the two types can be differentiated from each other by immunohistochemical methods.

Material and methods

HISTOLOGICAL SPECIMENS

Corneal buttons obtained at the time of penetrating keratoplasty from two Finnish patients with LD type I and three patients with LD type II were taken from the files of the ophthalmic pathology laboratory, Helsinki University Central Hospital, and three buttons from two American patients with LD type I from the department of ophthalmic pathology, Armed Forces Institute of Pathology (Table 1). Four eyes from three Finnish patients with LD type II were obtained at autopsy from the department of pathology, University of Helsinki.

All six patients with LD type II had typical clinical findings of familial amyloidosis, Finnish type (FAD), and a positive family history of the disease, and they came from the Kymenlaakso region in south eastern Finland, known to have a high prevalence of Meretoja’s syndrome. In the four patients with LD type I, the disease appeared to be an isolated phenomenon unrelated to symptoms of systemic amyloidosis. For control purposes, identically processed corneal buttons from patients with granular dystrophy (ages 38 and 43 years), keratoconus (ages 27, 44, and 50 years), and leukemia (ages 78 and 81 years) were selected.

All specimens were formalin fixed and paraffin embedded. Routine stains included haematoxylin and eosin, van Gieson, Congo red, Masson trichrome, and periodic acid Schiff. Criteria for identifying an amyloid deposit were birefringence and apple green dichroism observed under polarised optics after Congo red staining.
Table 1  Immunoreactivity of dystrophic deposits in corneal lattice and granular dystrophies with antibodies to amyloid P component, mutated gelsolin, and native human gelsolin

<table>
<thead>
<tr>
<th>Case</th>
<th>Sex</th>
<th>Age (years)</th>
<th>Diagnosis</th>
<th>Nationality</th>
<th>anti-AP (^a)</th>
<th>anti-FAF (^b)</th>
<th>GS-2C4 (^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>L1a</td>
<td>Male(^a)</td>
<td>21</td>
<td>Lattice dystrophy, type I</td>
<td>American</td>
<td>++</td>
<td>+</td>
<td>NA</td>
</tr>
<tr>
<td>L1b</td>
<td>Male(^a)</td>
<td>22</td>
<td>Lattice dystrophy, type I</td>
<td>American</td>
<td>+</td>
<td>+</td>
<td>NA</td>
</tr>
<tr>
<td>L2</td>
<td>Female(^a)</td>
<td>24</td>
<td>Lattice dystrophy, type I</td>
<td>American</td>
<td>++</td>
<td>+</td>
<td>NA</td>
</tr>
<tr>
<td>L3</td>
<td>Female</td>
<td>55</td>
<td>Lattice dystrophy, type I</td>
<td>Finnish</td>
<td>++</td>
<td>-</td>
<td>(+)</td>
</tr>
<tr>
<td>L4</td>
<td>Male</td>
<td>75</td>
<td>Lattice dystrophy, type I</td>
<td>Finnish</td>
<td>++</td>
<td>-</td>
<td>(+)</td>
</tr>
<tr>
<td>M1</td>
<td>Female</td>
<td>65</td>
<td>Lattice dystrophy, type II</td>
<td>Finnish</td>
<td>++</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>M2</td>
<td>Female</td>
<td>66</td>
<td>Lattice dystrophy, type II</td>
<td>Finnish</td>
<td>++</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>M3</td>
<td>Male</td>
<td>72</td>
<td>Lattice dystrophy, type II</td>
<td>Finnish</td>
<td>++</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>M4</td>
<td>Male</td>
<td>72</td>
<td>Lattice dystrophy, type II</td>
<td>Finnish</td>
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<td>+</td>
<td>+</td>
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<tr>
<td>M5</td>
<td>Female</td>
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<td>++</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>M6</td>
<td>Female</td>
<td>80</td>
<td>Lattice dystrophy, type II</td>
<td>Finnish</td>
<td>++</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>
| C1   | Female | 38 | Granular dystrophy | Finnish | - | - | -/+
| C2   | Female | 43 | Granular dystrophy | Finnish | - | - | -/+

\(^a\) AP amyloid P component; FAF=mutated variant gelsolin; GS=native gelsolin; ++=strong; + moderate; (±)=weak immunoreaction; NA=not applicable.

\(^b\) Both specimens from the same patient.

\(^c\) Focal reaction mainly in the periphery of the deposits.

\(^d\) Autopsy material.

\(*\) Keratoconus in addition to lattice dystrophy.

**Results**

**LD TYPE I**

By light microscopy, three specimens represented moderately advanced dystrophy (Fig 1A-D) with relatively scarce lattice lines within the corneal stroma in Congo red staining (Fig 1A), and two came from patients with advanced lattice dystrophy (Fig 1E-H) displaying many lattice lines of various sizes throughout the stroma (Fig 1E, F). A discontinuous layer of amyloid was present under Bowman's layer (Fig 1A), partly replacing it in the two advanced cases (Fig 1E, F).

The anti-AP antisera labelled diffusely lattice lines in all specimens studied (Fig 1B, G; Table 1). In four cases, it also reacted focally with amyloid deposits associated with Bowman's layer (Fig 1B, G). The epithelial basement membrane was not labelled. The anti-FAF antisera could label weakly the cytoplasm of corneal epithelial and endothelial cells, consistent with its weak cross reactivity with native gelsolin (Fig 1C, H). In two specimens, scattered stromal keratocytes were also labelled (Fig 1H). Amyloid deposits around Bowman's layer remained negative (Fig 1C, H). In two specimens, lattice lines were negative, but in three others the antisera faintly bound to their periphery (Fig 1C, H).

For technical reasons, two specimens remained totally negative with MAb GS-2C4 and could not be evaluated. It labelled the cytoplasm of corneal epithelial cells and endothelial cells, and a subpopulation of stromal keratocytes in the remaining specimens. It two of these specimens, focal immunoreaction was seen in the periphery of some lattice lines (Fig 1D).

**LD TYPE II**

By light microscopy, all six specimens contained relatively few and thin lattice lines (Fig 2A-E), which concentrated to the anterior and middle stroma. They were seen with equal frequency in the central and peripheral cornea. A variably thick, almost continuous deposition of amyloid was always present under Bowman's layer, and a thin line of amyloid was seen at the level of the epithelial basement membrane in some specimens (Fig 2A, B). Bowman's layer was often

**IMMUNOHISTOCHEMISTRY**

Immunoperoxidase staining was carried out using commercial versions (Vectastain ABC Elite kits for rabbit and mouse IgG; Vector Laboratories, Burlingame, CA, USA) of the avidin biotinylated peroxidase complex (ABC) method as has been described in detail.\(^{33}\)

A rabbit antiserum was raised against purified amyloid subunit obtained by gel filtration of amyloid fibrils isolated from kidneys of a patient suffering from FAF.\(^{21-23}\) This fraction is homologous to that of human gelsolin, starting at position 173, but it shows an amino acid substitution, asparagine for aspartic acid, at position 187.\(^{21}\) On immunoblot analysis, the antisera reacts strongly with the FAF amyloid subunit and weakly with normal gelsolin.\(^{22}\) It labels amyloid deposits in renal glomeruli, conjunctiva, skin, eccrine sweat glands, perineural sheaths, and blood vessel walls in FAF, but does not label these tissues in controls.\(^{20-23}\) Absorption with purified FAF amyloid subunit abolishes the immunoreaction.\(^{21}\) Cross reactivity has not been detected in several other types of amyloidosis.\(^{33}\) The optimal dilution in this study was 1:800.

Mouse monoclonal antibodies to human gelsolin (GS-2C4, IgG\(_1\), Lot 041H-4847; dilution 1:750) were obtained from Sigma (St Louis, MO, USA). It detects an epitope on a 47 kDa chymotryptic cleavage peptide corresponding to residues 407–755 of the 93 kDa plasma gelsolin and containing its carboxy terminal actin binding site.\(^{34}\) It does not react with the FAF amyloid protein, derived from the amino terminal end of gelsolin.\(^{21}\) A rabbit antiserum to human amyloid P component (A302, Lot 118; dilution 1:500) was purchased from Dakopatts (Glostrup, Denmark).

**CONTROL EXPERIMENTS**

Omitting the primary or secondary antibody or the ABC complex abolished the immunoreaction. Normal rabbit serum (Ortho Diagnostics, Stillwater, MN, USA) and an unrelated murine IgG\(_1\) antibody (anti-synaptophysin, SY 38; Boehringer Mannheim, Mannheim, Germany) were used as negative controls. Purified amyloid protein was no longer available for blocking.
discontinuous, and scar tissue with occasional amyloid deposits invaded the subepithelial space. Thin amyloid streaks were also seen between superficial corneal lamellae, particularly in the limbal region (Fig 2F).

The anti-AP antiserum labelled lattice lines, amyloid deposits under Bowman's layer, and amyloid streaks between corneal lamellae with a weak to moderate intensity (Fig 2C, G; Table 1). Immunoreaction was also seen focally at the level of epithelial basement membrane (Fig 2C) and in amyloid deposits invadig the subepithelial scar tissue. The anti-FAF antiserum strongly and consistently labelled lattice lines and amyloid deposits around Bowman's layer (Fig 2D). This immunoreaction was most extensive under the Bowman's layer, but was also often seen at the level of the epithelial basement membrane (Fig 2D), and could involve the entire Bowman's layer. Immunopositive streaks between corneal lamellae were seen especially near the limbal region (Fig 2H). MAb GS-2C4 gave a faint to moderate, non-homogeneous granular immunoreaction in many lattice lines, amyloid deposits under Bowman's layer, and in streaks of amyloid between corneal lamellae (Fig 2I). Other deposits

Figure 1  Immunohistochemistry of lattice corneal dystrophy type I (A, E, F Congo red; B-D, G, H immunoperoxidase staining). In moderately advanced dystrophy (A-D, case L3) Congo red (A; ×225) and antiserum to amyloid P component (AP) (B; ×250) reveal amyloid in lattice lines (a) and as discontinuous deposits (arrowhead) under Bowman's layer (b), whereas the corneal epithelium (ep) remains unreactive. The anti-FAF antiserum binds to desquamating corneal epithelial cells and labels faintly other epithelial (ep) layers, but lattice lines (a) and amyloid (arrowhead) under Bowman's layer (b) do not react with it (G; ×250). MAb GS-2C4 to gelsolin reacts strongly with the corneal epithelium (ep) and in a focal pattern with lattice lines (a), but it is not immunoreactive with amyloid deposits (arrowhead) under Bowman's layer (b) (D; ×250). Advanced lattice dystrophy (E-H, case L18) with abundant lattice lines (a), extending up to the endothelium (double arrowhead), and anterior deposits totally replacing Bowman's layer (arrowhead) are seen in Congo red stainings without (E; ×185) and with polarisation (F; ×185), as well as with the antiserum to AP (G; ×260). In addition to faint epithelial labelling (ep), the antiserum to FAF reacts with the periphery of lattice lines (a) and with single stromal keratocytes (double arrowhead), but it does not label amyloid deposits (arrowhead) under Bowman's layer or the core of lattice lines (H; ×330).
remained totally negative, however (Fig 2E). As in other specimens, it labelled the cytoplasm of corneal epithelial cells, along with scattered keratocytes, preferably in the anterior and middle stromal layers (Fig 2E).

**CONTROL MATERIAL.**

The amyloid deposits in LD types I and II did not react with normal rabbit serum or the control MAb of IgG, isotype. Amyloid was not found in any control specimen with Congo red or the anti-AP antiserum (Fig 3A, B). The anti-FAF antiserum labelled single desquamating corneal epithelial cells and, in a few specimens, scattered keratocytes, similar to those in some specimens of lattice dystrophy (Fig 1C, H). It faintly bound to the periphery of several deposits in granular dystrophy, resembling edge artefact (Fig 3C; Table 1). A strong reaction in many subepithelial deposits of granular dystrophy was seen with MAb GS-2C4 to gelsolin (Fig 3D). Otherwise, it bound to the corneal epithelium, endothelium, and scattered keratocytes.

**Discussion**

Traditionally, the lattice lines in LD type I tend to

![Figure 2](http://bjo.bmj.com/)

**Figure 2** Immunohistochemistry of lattice corneal dystrophy type II (A, B, F Congo red; C-E, G-I immunoperoxidase staining). Congo red staining (A-E, case M6) without (A; ×240) and with (B; ×240) polarisation reveals a thin layer of amyloid (double arrowhead) under Bowman’s layer, and stromal lattice lines (a). The epithelial (ep) basement membrane (arrowhead) is also labelled. Antisera to amyloid P component (AP) (C; ×300) and FAF (D; ×300) react with lattice lines (a), anterior stromal amyloid deposits (double arrowhead), and occasional streaks of amyloid between stromal lamellae, whereas only the latter labels the epithelial basement membrane (arrowhead). MAb GS-2C4 to gelsolin reacts with the cytoplasm of corneal epithelial cells (ep) and occasional stromal keratocytes (arrowheads), but in this specimen does not label the anterior stromal deposits (double arrowhead) or streaks between corneal lamellae (E; ×300). In peripheral cornea (F-I, case M5), abundant thin amyloid deposits (arrowheads) are observed between corneal lamellae, more so towards the limbus (lim) than the centre (cen) of the cornea (F; ×130). Amyloid deposits under Bowman’s layer (b) are also seen. The amyloid streaks (arrowheads) react moderately for AP (G; ×300), strongly for FAF (H; ×300), and weakly for gelsolin (I; ×300). The epithelium has detached from this autopsy specimen.
to be more numerous and thinner, they extend more deeply into the corneal stroma, and spare more of its periphery than do those in LD type II.\textsuperscript{12} The former becomes symptomatic earlier and progresses more rapidly than the latter, which rarely requires corneal transplantation, even though it becomes manifest by the age of 20.\textsuperscript{9} The clinical clue to LD type II is the typical facies with droopy eyelids and protruding lips, due to cutaneous and cranial nerve involvement by amyloid deposits.\textsuperscript{10–13}

In LD type II, an even layer of amyloid is seen at the level of the epithelial basement membrane and beneath Bowman’s layer, which may show breaks in continuity, and stromal lattice lines are thin.\textsuperscript{14–17} We are aware of only one previous paper that has specifically compared the histopathology of LD types I and II.\textsuperscript{18} It noted more frequent and coarse amyloid deposition between the epithelium and fragmented Bowman’s layer, thicker and more irregular amyloid deposits under Bowman’s layer, and more widespread distribution of lattice lines in LD type I.\textsuperscript{19} These general findings were confirmed in our material. Moreover, thin streaks of amyloid between corneal lamellae were noted in LD type II, particularly in the limbal region, possibly associated with stromal kerocytes, a feature not emphasised before.

The antiserum to the variant gelsolin amyloid subunit of a patient with Meretoja’s syndrome always labelled lattice lines, as well as amyloid deposits on both sides of Bowman’s layer and between corneal lamellae in LD type II, in line with preliminary observations in one American and three Finnish patients with FAF.\textsuperscript{20,21} Immunoblotting has shown that the mutated gelsolin polypeptide is indeed present in the affected cornea.\textsuperscript{22} MAb GS-2C4 to native human gelsolin\textsuperscript{23} reacted focally with corneal amyloid deposits in LD type II, consistent with the fact that, in many types of amyloidosis, variable amounts of the precursor protein can be deposited along with the amyloid polypeptide. Indeed, in some patients with FAF at least, the normal allele may be coexpressed with the mutated one.\textsuperscript{24} Since MAb GS-2C4 recognises a carboxyl terminal epitope of gelsolin,\textsuperscript{25} absent from the FAF amyloid fragment derived from its amino terminal end, it is not a very useful reagent for demonstration of LD type II. Identical reactivity has nevertheless been reported in two previously studied specimens.\textsuperscript{25–27}

Convincing labelling of amyloid deposits was not obtained with the anti-FAF antiserum in LD type I. MAb GS-2C4 to gelsolin was initially reported to be non-reactive with LD type I and other corneal amyloidoses.\textsuperscript{28} More recently, a moderate to intense immunoreaction, mainly bordering lattice lines, was reported in four of six corneal buttons in LD type I.\textsuperscript{29} Similar immunoreaction was occasionally seen in polymorphic amyloid degeneration, primary familial subepithelial amyloidosis, and secondary corneal amyloidosis, and it was speculated to result from changes in molecular conformation of presumed abnormal gelsolin fragments in the centre of these deposits.\textsuperscript{30} In the present study, not only two of three evaluable specimens of LD type I, but also both cases of granular dystrophy reacted focally with MAb GS-2C4, suggesting that plasma gelsolin may bind to several types of corneal deposit. Thus, the significance of focal binding of MAb GS-2C4 to amyloid deposits other than LD type II remains open to discussion.

These results are insufficient to prove that LD type I is unrelated to gelsolin, as it might result from another mutation in the gelsolin gene or a different degradation pathway of the amyloid precursor. They are, nevertheless, consistent with a recent linkage analysis in a large kindred, which seemed to exclude gelsolin as a candidate gene for LD type I in this particular family.\textsuperscript{31} Failure to label the amyloid deposits with anti-

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Figure 3 Immunohistochemistry of control specimens with granular dystrophy (A Masson trichrome; B, D immunoperoxidase staining). Light microscopy (A–D, case C2) reveals dystrophic deposits both deep within the stroma (g) and subepithelially (arrowhead) (A, ×150). Neither anterior (arrowhead) nor deep deposits (g) react reliably with the anti-amyloid P component (B, ×150) and anti-FAF antiserum (C, ×300), respectively, giving edge artefact-like reaction, but MAb GS-2C4 to gelsolin labels quite strongly the subepithelial deposits (arrowheads) (D, ×150).
bodies to gelsolin was reported in this kind. However, because LD type I is quite a heterogeneous disease as to its clinical presentation, age of onset, laterality, and degree of stromal involvement, it is not yet certain that the precursor protein is identical in all families.

Finally, amyloid deposits in LD type I and II reacted for amyloid P component (AP), a plasma protein probably deposited along with the amyloidogenic protein in all types of amyloidosis. Whereas most, but not all, pathologic studies have confirmed its presence in LD types II517 and III, highly conflicting results have been reported for LD type I. The presence of AP in all types of LD is perhaps to be expected, and it cannot be used to differentiate between them.

Although molecular genetic studies are likely to become the most reliable means of differentiating between various types of lattice dystrophy in the clinical setting, and identifying affected family members before they develop biomicroscopically detectable corneal changes, the amyloid technique, the antiserum against gelsolin, and the presence of AP in the lattice corneal specimens to determine whether there is evidence of LD type II. As the antigen is resistant to formalin fixation and paraffin embedding, retrospective studies of old corneal buttons will be possible. Meretoja's syndrome is a relatively unknown and most probably underdiagnosed condition, and such studies might lead to a better understanding of its world distribution and epidemiology.

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