The APO*E3-Leiden mouse as an animal model for basal laminar deposit

Mike Kliffen, Esther Lutgens, Mat J A P Daemen, Ebo D de Muinck, Cornelia M Mooy, Paulus T V M de Jong

Abstract

Aim—To investigate the APO*E3-Leiden mouse as an animal model for age related maculopathy (ARM) related extracellular deposits.

Methods—Eyes were obtained from APO*E3-Leiden transgenic mice on a high fat/cholesterol (HFC) diet (n=12) or on a normal mouse chow (n=6), for 9 months. As controls, eyes were collected from APO-E knockout mice on the same diets. From each mouse one eye was processed for microscopic evaluation and immunohistochemistry with a polyclonal antibody directed against human apo-E. Electron microscopy was also performed.

Results—All 12 eyes of the APO*E3-Leiden mice on an HFC diet contained basal laminar deposit (BLD; class 1 to class 3), whereas two of six APO*E3-Leiden mice on normal chow showed BLD class 1. The ultrastructural aspects of this BLD were comparable with those seen in early BLD in humans, and BLD showed immunoreaction with anti-human-apo-E antibodies. No BLD was found in any of the control mice. Drusen were not detected in any of the mice.

Conclusion—These results indicate that APO*E3-Leiden mice can be used as animal model for the pathogenesis of BLD, and that a HFC diet enhances the accumulation of this deposit. Furthermore, this study supports the previously suggested involvement of dysfunctional apo-E in the accumulation of extracellular deposits in ARM.

Materials and methods

MICE
All procedures complied with the Association for Research in Vision and Ophthalmology statement for the use of animals in ophthalmic and vision research.

Transgenic mice expressing the human APO*E3-Leiden gene (and producing a dysfunctional form of human apo-E3), and APO-E knockout mice were generated as described earlier.16 17 For the present study, mice of the F10 generation were used. Before entry into the study, animals were kept on a standard mouse chow (SRM-A, Hope Farms, Woerden, Netherlands). At the mean age of 37 (SD 0.6) weeks, 12 APO*E3-Leiden mice and 12 APO-E knockout mice were put on a high fat/cholesterol (HFC) diet containing 15% cacao butter, 0.5% cholate, 1% cholesterol, 40.5% sucrose, 10% corn starch, 1% corn oil, and 4.7% cellulose (Hope Farms) for 9 months. Six APO*E3-Leiden mice and 12 APO-E knockout mice received standard mouse chow for 9 months.

The mice in this study were primarily used for research on atherosclerosis.18 All mice were housed under standard conditions on the shelf below the top one in a room with normal fluorescent light exposure. After completion of the
diet according to protocol, each mouse was anaesthetised with xylazine (0.0025 ml/g) and ketamine (0.001 ml/g) intraperitoneally, and the arterial tree was perfused with 0.9% NaCl (3 minutes) and 10% phosphate buffered formalin (pH 7.4, 3 minutes), both containing 20% nitroglycerine. Subsequently, both eyes were taken out and fixed overnight in 10% phosphate buffered formalin. One of the eyes (random) of each mouse was used for this study and the central retina was embedded in paraffin for histopathological classification and immunohistochemistry.

**HISTOPATHOLOGICAL CLASSIFICATION OF MOUSE CENTRAL RETINA**

Paraffin sections (5 µm) were stained with haematoxylin and eosin, the periodic acid Schiff reaction, and Mallory staining. The retinas were histologically classified according to the independent grading of both drusen and BLD. This classification of drusen and BLD has been described earlier in human eyes. In brief: no BLD (class 0); small solitary patches of BLD (class 1); a thin continuous layer of BLD (class 2); a thick layer of BLD of at least half the height of the RPE cells (class 3). No drusen (class 0); 1–3 drusen (class 1); 4–10 drusen (class 2); many or confluent drusen (class 3).

**IMMUNOHISTOCHEMISTRY**

Paraffin sections (5 µm) of all retinas were mounted on 3-aminopropyl-tri-oxi-silane (Sigma, St Louis, MO, USA) coated glass slides. After dewaxing and rehydration they were rinsed with water and phosphate buffered saline, and incubated with 5.5 mU/ml pronase E (Sigma, St Louis, MO, USA) for 6 minutes at 37°C. The slides were then placed in a Sequenza Immunostaining Workstation (Life Sciences International, Velthoven, Netherlands), and incubated for 30 minutes with a rabbit polyclonal antibody directed against human apo-E (kindly supplied by L Havekes from the Gaubius Laboratory, Leiden, Netherlands). This antibody does not cross react with mouse apo-E. After washing with phosphate buffered saline, the slides were incubated for 30 minutes with biotinylated secondary antibodies (Multilink, 1:75 dilution, Biogenex, San Ramon, MO, USA). The slides were washed again and incubated for 30 minutes with alkaline phosphatase conjugated streptavidin (Multilink, 1:75 dilution, Biogenex, San Ramon, MO, USA) coated glass slides. After dewaxing and rehydration they were mounted on uncoated mesh 300 copper grids. Ultrathin sections (70–80 nm thickness) were mounted on uncoated mesh 300 copper grids. After staining for 30 minutes with uranyl acetate and 2 minutes with lead citrate, the sections were examined with a Zeiss EM 902 transmission electron microscope.

**RESULTS**

**HISTOPATHOLOGICAL CLASSIFICATION OF MOUSE CENTRAL RETINA**

All 12 eyes of the APO*E3-Leiden mice on a HFC diet for 9 months contained BLD. Five eyes had BLD class 1, six eyes had BLD class 2, and one eye had BLD class 3 (Fig 1A). Of the six APO*E3-Leiden mice fed a normal mouse chow, two eyes contained BLD class 1, three eyes had no BLD (Fig 1B), and one eye could not be investigated properly because of overlapping tissue layers after processing.

None of the APO-E knockout mice (either on HFC or normal diet) contained any BLD (that is, class 0) or any type of drusen (Fig 1C and D). There were no differences between the retinas of APO*E3-Leiden mice and APO-E knockout mice by light microscopy.

None of the eyes had any type of drusen.

**IMMUNOHISTOCHEMICAL LOCALISATION OF APOLIPOPROTEIN-E**

Human apo-E was immunohistochemically localised in BLD in all eyes of APO*E3-Leiden mice containing BLD (Fig 1E). In APO*E3-Leiden mice a diffuse staining was noted throughout the BLD and over some plasma remnants in blood vessels.

No positive staining with the apo-E antibody in APO-E knockout mice was found.

**ELECTRON MICROSCOPY**

Utrastructurally, the major part of BLD (located between the RPE cell membrane and its basement membrane) in these APO*E3-Leiden mice appeared as homogeneous basement membrane-like material (Fig 2), but also minute amounts of fibrous long spacing collagen were present (not shown).

**Discussion**

We have demonstrated the presence of BLD in all eyes from APO*E3-Leiden mice on a HFC diet for 9 months. Additionally, 33% of eyes from APO*E3-Leiden mice on a normal mouse chow also showed BLD. This mouse BLD showed positive immunoreactivity for human apo-E. Because the human apo-E antibody does not cross react with mouse apo-E, and APO-E knockout mice did not show positive immunoreactivity at all, it is reasonable to assume that the immunoreactivity of BLD in APO*E3-Leiden mice was solely due to the presence of human apo-E.

Various ratios between the individual components of human BLD have been described. The ultrastructural aspects of the mouse BLD we have found in this study are comparable with early BLD in humans.
Vesicular deposits such as those described in diffuse thickening of Bruch’s membrane (sometimes referred to as basal linear deposit) were not seen in this BLD.4

It has been reported recently that wild type C57BL/6 mice on an HFC diet do not develop light microscopically detectable amounts of extracellular deposits, although some diffuse thickening of Bruch’s membrane was demonstrated with electron microscopy.25 The eyes used in this study were obtained from mice that were used in an atherosclerosis related research project. The perfusion fixation technique (based on formaldehyde) that was used in this study resulted in optimal eye tissue preservation for (immuno)histochemical purposes. However, this technique is not optimal for electron microscopy, and could have created some artefacts. Therefore, we only used electron microscopy for the location and ultrastructure of relatively large amounts of BLD, and not for possible minute changes not detectable by light microscopy (for example, in Bruch’s membrane). Further studies especially designed for electron microscopy need to be done in order to elucidate the changes in RPE and Bruch’s membrane in these mice.

Recently, we have demonstrated apo-E in BLD and drusen in human maculas, and we have found a decreased risk for AMD in people carrying an apo-e4 allele compared with people homozygous for apo-e3.16 Most animals, including mice, carry only one allele of apo-E that is E4-like.24 This allele is much more efficient in its function (for example, in its receptor binding properties) than apo-E3.27 It is thought that this is one of the reasons that normal mice are very resistant to the development of atherosclerosis.28 It has been demonstrated previously that APO*E3-Leiden and APO-E knockout mice on an HFC diet develop severe atherosclerosis,29 with many features that are characteristic of human atherosclerotic plaques.25 Therefore, we have chosen the APO-E knockout mice as a control group instead of normal (apo-E4 carrying) mice, which would not have developed this severe atherosclerosis. APO*E3-Leiden and APO-E knockout mice on normal mouse chow do not develop atherosclerosis. The fact that more BLD was found in this model in atherosclerotic APO*E3-Leiden animals than in non-atherosclerotic APO*E3-Leiden animals supports the association between AMD and atherosclerosis that has been found in human epidemiologic studies.30 On the other hand APO-E knockout mice on an HFC diet do develop atherosclerosis but no BLD. Therefore, it seems likely that BLD in these APO*E3-Leiden mice is related to a dysfunctional apo-e, and that this BLD is attenuated by atherosclerosis/hyperlipidaemia.

Drusen and Bruch’s membrane contain various amounts of phospholipids and neutral lipids. These lipids are assumed to cause a diffusion barrier between the choriocapillaris and the RPE, but the mechanism by which these lipids accumulate is unknown.11–13 Most research points to accumulation of inappropriately metabolised products from the RPE, but
also serum derived origin should be considered. Since APO*E3-Leiden has a lower affinity for the LDL receptor, and causes a higher plasma level of apo-E, APO*E3-Leiden in BLD could be derived from the choriocapillaris. On the other hand it could be hypothesised that apo-E plays a part in the transport of retinal lipids, which have a high turnover, especially in the photoreceptors. A dysfunctional APO*E3-Leiden could alter the lipid transport system, and cause accumulation of BLD. Degeneration of photoreceptors is a main event in the end stages of ARM. Therefore, the presence of apo-E in BLD could be due to the known involvement of apo-E in the repair of damaged neural tissue. Recently, Schneeberger et al demonstrated the presence of apo-E in the subretinal space in several forms of exudative retinal detachments, which could support this hypothesis. Furthermore, apo-E also accumulates in extracellular deposits in Alzheimer’s disease, another neurodegenerative disease. Another explanation for the presence of apo-E in BLD could be the fact that apo-E is easily bound to glycosaminoglycans, molecules that are known to be present in BLD. One of the major problems in studying ARM is the lack of fresh human tissue necessary for most of the biochemical techniques. Furthermore, the macular area in the eye consists of only a small amount of tissue. A piece of macular tissue of one eye (about 0.1 cm²) contains about 2 mg total protein. Highly specialized techniques—for example, two dimensional polyacrylamide gel electrophoresis and two dimensional electrophoresis on cellulose acetate membranes, have to be used to cope with these small amounts of human eye tissue. Therefore, the APO*E3-Leiden mouse model could prove to be helpful for further investigation (on the origin) of BLD. It also provides an opportunity to collect BLD within a few minutes post mortem, which is necessary for many biochemical techniques, but almost impossible from a human source. Additionally, this animal model could be used, despite the lack of a real macula, in intervention studies in order to prevent the accumulation of BLD, which is histologically an important feature associated with the development of ARM.

In conclusion our results indicate that APO*E3-Leiden mice can be used as an animal model for the pathogenesis of BLD, and that an HFC diet enhances the accumulation of this deposit. Furthermore, this study supports the previously suggested involvement of dysfunctional apo-E in the accumulation of extracellular deposits in ARM.

19 Van Ree JH, van den Broek WJ, van der Zee A, et al. Inactivation of Apo-e and Apo-e1 by two consecutive rounds of

www.bjophthalmol.com


