ApoB100,LDLR−/− Mice Exhibit Reduced Electroretinographic Response and Cholesteryl Esters Deposits in the Retina

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PURPOSE. To evaluate the retinal phenotype of 7- and 14-month-old apoB100,LDLR−/− mice, a relevant animal model of lipid metabolism dysfunction.

METHODS. Single-flash electroretinograms were obtained from 7- and 14-month-old apoB100,LDLR−/− and control mice fed a standard diet under both scotopic and photopic conditions. Visual cycle retinoids were analyzed in eyes from dark-adapted mice. Retinal and choroidal vascularization was evaluated with scanning laser ophthalmoscopy. Fatty acids were analyzed in the retina. Esterified and free cholesterol was detected in eye cryosections.

RESULTS. Scotopic and photopic b-wave amplitudes were significantly reduced in apoB100,LDLR−/− mice compared with control mice at 7 and 14 months of age (between -25% and -35% in 7-month-old animals and between -50% and -60% in 14-month-old animals at 25 cds/m2). Esterified cholesterol was found to accumulate at the basement of the retinal pigment epithelium in apoB100,LDLR−/− mouse eyes. On the contrary, no significant changes in the retinal profile of fatty acids and visual retinoids were observed in apoB100,LDLR−/− mice compared with control animals.

CONCLUSIONS. The exclusive expression of apoB100 in LDL receptor-null mouse altered the ERG profile, without modifying the visual cycle of retinoids and led to cholesterol deposition in the retina. These findings clearly suggest the role of cholesterol metabolism in the functioning of the retina and possibly in the etiology of ocular diseases, including age-related macular degeneration. (Invest Ophthalmol Vis Sci. 2008;49: 1307–1314) DOI:10.1167/iovs.07-0808

Accumulation of lipids under the retinal pigment epithelium (RPE) and within Bruch's membrane (BrM) is a normal feature of aging1,2 and has also been observed in human eyes with age-related maculopathy (ARM).2,5 The mechanism of lipid deposition in sub-RPE is partly unknown but several hypotheses have emerged. Plasma lipids may infiltrate BrM.4 Alternatively, lipid deposits may originate from abnormal RPE lipid metabolism involving apolipoprotein B (apoB)-rich lipoprotein synthesis.2,5

Insights have come from various mouse models of cardiovascular disease. C57Bl6 mice fed a high-fat diet and exposed to blue-green light develop basal laminar deposits,6 characterized by unesterified cholesterol accumulation,7 a histologic feature found in patients with ARM8 or age-related macular degeneration (AMD).8 The role of the low-density lipoprotein receptor (LDLR) has recently been highlighted in the transgenic LDLR knock out (LDLR−/−) mouse model. After being fed a high-fat diet, LDLR−/− mice exhibited BrM abnormalities characterized by BrM thickening and deposition of translucent material9 that may be lipid-rich particles comparable to the basal laminar deposits and drusen observed in patients with AMD.10 The relative resistance of these mice to development of atherosclerotic lesions, unless they consume a high-cholesterol atherogenic diet,11 has been attributed to the clearence of apoB48-lipoproteins by receptors other than LDLR, leading to lower plasma levels of LDL than can be expected from the lack of interaction between LDLR and apoB100, and subsequent lipoprotein clearance. Mice expressing exclusively apoB100 showed a modest increase in plasma LDL cholesterol but developed atherosclerosis12,13 and basal laminar deposits in BrM14 only after ingestion of an atherogenic diet. These features limit the usefulness of the LDLR−/− and apoB100 transgenic mice for studies on the role of LDLR and apoB100 in the context of an atherogenic diet, but fail to analyze the endogenous effect of pure LDL clearance deficiency independent of the increase in plasma lipid induced by the diet.

Transgenic mice lacking LDLR and expressing exclusively apoB100 have been established as a valuable model for atherosclerosis on a low-fat diet, with high plasma cholesterol levels and aortic atherosclerotic plaques.15 These mice offer a unique opportunity to test the hypothesis that the lack of apoB100 and LDLR interaction alters lipid metabolism and impairs functioning of the retina. Therefore, we sought to analyze the electroretinographic profile, fundus images, and retinal and choroidal vasculature in 7- and 14-month-old apoB100,LDLR−/− mice, by comparison with control mice. In parallel, esterified and unesterified cholesterol deposition was evaluated in eye sections using the fluorescent probe filipin.

MATERIALS AND METHODS

Mice

The use of animals was in accordance with the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research. French legal and institutional ethics committee review board approvals were obtained.

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B6129SF2/J and transgenic apoB100,LDLR−/− mice were purchased from Jackson Laboratories (Bar Harbor, ME), mated, bred, and maintained under controlled conditions of temperature (22 ± 1°C), hygrometry (55%–60%), and light (lights on 7 AM–7 PM, 18 lux) until the age of 7 and 14 months (animal breeding house; INRA, Dijon, France). The mice were fed a standard low-fat diet (128S; Harlan-Teklad, Gannat, France).

Electroretinographic Evaluation

Before ERG recording and fundus examination, the mice were anesthetized by SC injection of ketamine (66.7 mg/kg body weight; Ketanest; Parke-Davis, Berlin, Germany) and xylazine (11.7 mg/kg body weight; Rompun; Bayer Vital, Leverkusen, Germany), and their pupils were dilated with tropicamide eye drops (Mydracium Stullin; Pharma Stullin, Nabburg, Germany).

Fundus Photography

In vivo black-and-white images of the fundus were obtained by angiography (Heidelberg Retina Angiograph [HRA]; Heidelberg Engineering GmbH, Heidelberg, Germany), as described elsewhere.16 The HRA is a confocal scanning laser ophthalmoscope (SLO) that uses different lasers deflected by a mirror oscillating sinusoidally at a frequency up to 20 Hz. Reflectance images of the fundus were acquired using the infrared diode laser (810 nm) and the argon green laser (514 nm). Variation of the focus of the laser beam allowed visualization of the different layers of the fundus (i.e., retina versus RPE). The size of the square scan field was set to 20° × 20° and 10° × 10°. Autofluorescence was excited by the argon blue wavelength (488 nm).

SLO Angiography

Retinal and choroidal vasculature changes were evaluated as described elsewhere.16 After an SC injection of 1% fluorescein (0.25 mL/30 g body weight) and 0.4% ICG (0.5 mL/30 g body weight). The retinal and choroidal vessels began filling approximately 30 seconds after administration. Eleven minutes after dye administration, pictures from the retinal and choroidal vasculature were recorded with two lasers of the angiography and infrared diode laser (795 nm) for excitation during ICG angiography. Barrier filters at 500 and 810 nm provided the stable level of the photopic response.18 Amplitudes as a function of flash intensity were fit with a nonlinear saturating function (the Naka-Rushton equation) to obtain values of Rmax (a maximum response amplitude parameter) and log σ (a sensitivity parameter that represents the flash intensity that produces one-half the maximum response amplitude).

Analysis of the Visual Cycle Retinoids

All procedures with retinoids, from the euthanizing of the animals to analysis, were performed under red illumination to avoid photosensorization and degradation of the retinoids. Fourteen-month-old apoB100,LDLR−/− and control mice were dark-adapted for 24 hours before death. The eyes were enucleated within 30 seconds after decapitation. Whole mouse eyes (six per analysis) were homogenized in 1.5 mL of 0.1 M MOPS (pH 6.5), 0.2% SDS, and 10 mM NH4OH in a glass–glass homogenizer and incubated for 30 minutes at room temperature to allow retinal oximes to form. Retinoids were then extracted twice with 4 mL hexane after the addition of 1 mL ethanol. After centrifugation at 900 g for 2 minutes, the upper phases were collected, dried under nitrogen gas, and resuspended in 150 μL hexane. Retinoids (50 μL) were injected onto a silica HPLC column (250 × 4.6 mm ID, 5 μm; Intersil Si; Varian, Les Ulis, France) equilibrated with hexane. Retinoids were resolved in hexane-ethyl acetate (90:10, vol/vol) at a flow rate of 1 mL/min. The absorbance of the effluent was monitored simultaneously at 325 nm (for retinols and retinyl esters) and 357 nm (for retinal oximes). Retinyl esters were separated with methanol on a reversed-phase HPLC column (250 × 4.6 mm ID, 5 μm; Nucleosil C-18; Varian) and detected at 325 nm. Retinoids were identified by comparison with commercial standards of retinyl laurate, retinyl myristate, retinyl palmitate, all-trans retinol, 13-cis-retinol and all-trans retinal (Sigma-Aldrich, St. Quentin Fallavier, France).

Eye Section Preparation

Seven- and 14-month-old mice were killed, blood samples were taken, and the eyes were enucleated and embedded in a cryoprotectant resin (Cryomount, Thermo-Shandon, Pittsburgh, PA). For further detection of unesterified (UC) and esterified (EC) cholesterol, the eyes were serially sectioned at 8 μm, dried overnight at room temperature, and stored at −80°C until used. For morphometric analyses, the eyes were sectioned sagittally at 20 μm so that the entire eye from the ora serrata to the optic nerve was visible. Sections were fixed 10 minutes in acetone and stored at −20°C until used.

Filipin Histochemistry for Detecting UC and EC

Cholesterol was revealed in eye sections (n = 4 in each strain of mice at each age, five sections per eye) with filipin,19,20 a fluorescent probe that specifically binds the 3β-hydroxy group of cholesterol, according to a protocol described by others.3,7 Sections for determination of UC were hydrated and incubated for 30 minutes in a filipin solution consisting of 5 mg of filipin (Sigma-Aldrich) dissolved in 1 mL of dimethylformamide and diluted in 100 mL of phosphate-buffered saline. Sections for determination of EC were processed as follows: Native UC was first extracted by two 5-minute rinses in 70% ethanol, and native EC was then hydrolyzed with cholesterol esterase (EC 3.1.1.13; Sigma-Aldrich) at a concentration of 1.65 U/mL in 0.1 M potassium phosphate buffer (pH 7.4), for 3 hours at 37°C. UC newly released by the hydrolysis of EC was then stained with filipin, as just described. Sections incubated with the enzyme vehicle served as the control. The sections were viewed by epifluorescence microscopy with a 420-nm excitation filter and 520-nm barrier filter.

Morphometric Analysis

To determine the thickness of the outer nuclear layer (ONL), we stained the sections with hematoxylin and eosin (Mayer Hematoxylin, Harris Gill Hematoxylin II PAP 1:1, vol/vol; Laborond, Templemars, France). The sections that contained the optic nerve were bright-field imaged with a microscope (Eclipse E400; Nikon, Champigny-sur-Marne, France) at 10× power and photographed with a 3 CCD color
video camera (KY-F55E, JVC; Victor Company of Japan, Tokyo, Japan). Digital images were analyzed on computer to measure the thickness of the different layers of the retina ($n = 3$ measurements taken between 0.300 and 0.700 mm from the optic nerve, from four eyes in each strain of animals; Optimas 6.5; IPS, North Reading, MA). Data were analyzed by a standard $t$-test procedure (SAS software; SAS Institute, Cary, NC). Differences were considered significant at $P < 0.05$.

**Fatty Acid Profile Analysis of the Retina**

At the ages of 7 and 14 months, the mice were euthanatized, the eyes enucleated, and the neural retina removed and stored in chloroform.

**FIGURE 1.** Electroretinographic data from 7-month-old control and apoB100,LDLR$^{-/-}$ mice. Dark-adapted (a) a- and (b) b-wave amplitudes and (c) light-adapted b-wave amplitude from apoB100,LDLR$^{-/-}$ mice are plotted as a function of the flash intensity. Red lines: normal range given by the 5% and 95% quantile of the control mice ($n = 6$).

**FIGURE 2.** Electroretinographic data from 14-month-old control and apoB100,LDLR$^{-/-}$ mice. Dark-adapted (a) a- and (b) b-wave amplitudes and light-adapted b-wave amplitude (c) from apoB100,LDLR$^{-/-}$ mice are plotted as a function of the flash intensity. Red lines: normal range given by the 5% and 95% quantile of the control mice ($n = 6$).
methanol (2:1, vol/vol). Lipids were extracted, and fatty acids were transmethylated according to Morrison and Smith. Fatty acid methyl esters were analyzed by gas chromatograph (5890 series II; Hewlett-Packard, Palo Alto, CA) equipped with a split/splitless injector, a flame-ionization detector, and a CPSil88-silica capillary column (100 m x 0.25 mm ID, film thickness 0.20 μm; Varian). The injector and the detector were maintained at 250°C and 280°C, respectively. Hydrogen was used as a carrier gas (inlet pressure, 210 kPa). The oven temperature was fixed at 60°C for 1 minute, increased to 85°C at a rate of 3°C/min and then to 190°C at a rate of 20°C/min and left at this temperature for 65 minutes. Fatty acid methyl esters were identified by comparison with commercial standards.

**Plasma Levels of Cholesterol**

Plasma was obtained from single animals after the collection of whole blood in heparin by aortic puncture at the time of euthanization. Total plasma cholesterol was measured by using an enzymatic procedure. Data were analyzed by a standard two-factor (age, strain) ANOVA procedure (SAS software; SAS Institute). Differences were considered significant at \( P < 0.05 \).

**RESULTS**

The apoB100,LDLR\(^{-/-}\) mouse serves as an established model for atherosclerotic mechanisms because of the elevation of plasma cholesterol, even in animals fed a standard low-fat diet. We first confirmed this finding, showing that plasma cholesterol was increased threefold in apoB100,LDLR\(^{-/-}\) mice compared with the control at 7 (3.3 ± 0.09 and 1.0 ± 0.33 mg/mL, respectively, \( n = 6 \) in each group; \( P < 0.0001 \)) and 14 months of age (3.7 ± 0.07 mg/mL and 1.2 ± 0.07 mg/mL, respectively, \( n = 5 \) in each group; \( P < 0.0001 \)).

**ERG Response**

ERGs were recorded from 7- and 14-month-old apoB100,LDLR\(^{-/-}\) mice under dark-adapted (Fig. 1a, 1b for 7-month-old mice, Fig. 2a, 2b for 14-month-old mice) and light-adapted (Fig. 1c for 7-month-old mice, Fig. 2c for 14-month-old mice) conditions. Scotopic latencies of both a- and b-waves were unchanged in 7- and 14-month-old apoB100,LDLR\(^{-/-}\) mice when compared with control mice at the corresponding age (data not shown). Similarly, under photopic conditions, no delay of b-wave latency was observed (data not shown). The amplitude of a- and b-wave was reduced as a function of time between 7 and 14 months in both strains of mice, and in apoB100,LDLR\(^{-/-}\) mice at both ages compared to age-matched control mice (compare Figs. 1a, 1b for 7-month-old mice, Fig. 2a, 2b). The Naka-Rushton representation of the b-wave amplitude as a function of stimulus intensity was used to calculate \( R_{\text{max}} \), as a parameter for the maximum-response amplitude, and \( \log \sigma \), the sensitivity parameter that represents the flash intensity that produces one half the maximum response amplitude.

**Fatty Acid Composition of the Neural Retina**

Fatty acid composition of the neural retina is of major concern for retinal function, as depicted in studies showing that docosahexaenoic acid (DHA) enhances the kinetics of the photocycle by interacting with rhodopsin and that DHA depletion lowers the electroretinographic response. Increased autofluorescence in a large number of perivascular cells (Fig. 4d) was present in all apoB100,LDLR\(^{-/-}\) mice but not in the control mice, whereas the vascular structure in the 514-nm surface image (Fig. 4e) and the fluorescein angiography (Fig. 4f) at 488 nm did not show any abnormalities.

**Detection of UC and EC in Eye Sections**

UC and EC were detected by using filipin in eye sections of both control and apoB100,LDLR\(^{-/-}\) mice at 7 and 14 months. UC appeared to be broadly distributed throughout the retina of both control and apoB100,LDLR\(^{-/-}\) mice (Figs. 5a, 5c, 5e, 5g). The most striking difference between apoB100,LDLR\(^{-/-}\) and
control mice was the age-related accumulation of EC in apoB100,LDLR–/– at 7 months and to a greater extent at 14 months (Figs. 5b versus 5d, 5f versus 5h). Intense laminar fluorescence of EC was found at the basement of the RPE (Figs. 5d, 5h, 5i). On the contrary, 7- and 14-month-old control mice displayed similar low EC staining (Figs. 5b, 5f).

**Morphometric Analysis of the Retina**

The mean thickness of the outer layers of the retina in apoB100,LDLR–/– mice was comparable to that in control animals (outer segments [OS]: 40.8 ± 7.9 μm compared with 33.5 ± 4.7 μm; ONL: 54.7 ± 8.1 μm compared with 57.1 ± 11.9 μm, respectively, in 14-month-old mice, P > 0.05).

**DISCUSSION**

This report describes a correlation between a dysregulation of the lipid metabolism triggered by the exclusive expression of the human apoB100 gene in the LDL knockout mouse and a reduced ERG response, indicating a functional impairment. As lipids are major components of the retina and account for up to 50% of rod outer segment membrane dry matter, a change in fatty acid status may be etiologically involved in the reduction of the ERG response. In particular, the interaction between rhodopsin and fatty acids may be relevant, since rhodopsin and DHA colocalize in the outer segments of photoreceptors. DHA is recognized as a key factor in the visual function by creating tight associations with rhodopsin that may facilitate the transition to its active form and enhance the kinetics of the photocycle.Investigators in numerous studies have reported reduced ERG-wave amplitude in DHA-depleted animals. A degeneration of rods and cones would lead to low content in DHA and rhodopsin in the neural retina and thinning of the outer layers of the retina (OS and ONL) that were not observed in our study. Consequently, our data showing reduced ERG in old apoB100,LDLR–/– mice cannot be explained by this type of DHA–rhodopsin mechanism, since no DHA depletion was reported in the retina of these animals, and no differences in the thickness of the OS and ONL were observed between control and apoB100,LDLR–/– mice. Similarly, the age-dependent reduction of b-wave amplitude in both control and apoB100,LDLR–/– mice does not correlate with a loss of DHA in the retina.

Hyperlipemia in the null mice may also contribute to the lower ERG response. Recent clinical data from a single patient showed that hyperlipemia was associated with lipemia retinalis and reduced ERG, although no signs of vascular abnormalities were observed. In this patient, the ERG response returned to normal when hyperlipemia was corrected. The authors hypothesized that the presence of large lipid-rich particles in the blood may produce a turbid serum appearance that would reduce the light intensity reaching the photoreceptors and consequently decrease the ERG response. This potential mechanism remains improbable in our model, since one would expect a- and b-wave latencies to increase concomitantly with the turbidity of the ocular fluids. However, no such delay was observed in apoB100,LDLR–/– mice (data not shown), which discredits the involvement of hyperlipemia, per se.

Far fewer data are available on the role of cholesterol in ERG. Cholesterol was found to form nonspecific interactions with rhodopsin in vitro, which we assume to have only a slight and inconsistent effect on the retinoid cycle. Indeed, like others, we reported only low content of free cholesterol in the outer segment of the neural retina, which renders the effect of any interaction between cholesterol and rhodopsin highly putative. Moreover, this mechanism remains improbable even in 14-month-old apoB100,LDLR–/– mice since, although cholesteryl ester deposition was observed at the basement of RPE in the null mice (Fig. 5), no increase in free cholesterol was found in the outer segments.

We presume the abnormal lipid metabolism in the retina of apoB100,LDLR–/– mice is involved in the accumulation of cholesteryl ester at the basement of RPE. The apoB100,LDLR–/– mouse is an established model for atherosclerosis, in that it shares close features with atherosclerosis in humans: enhanced plasma cholesterol concentrations, aortic fatty streaks, presence of LDL that is almost the exclusive form of cholesterol transport in plasma, and the exclusive expression of human apoB100. Apo-B100 is a large protein that is produced in the liver, where it is assembled into VLDL and remains present in LDL. A commonly recognized role for apoB100 is its capacity to bind LDL, thereby facilitating internalization of cholesteryl esters from LDL into peripheral cells. The expression of apoB100 in the retina of apoB100,LDLR–/– mice remains to be shown. Nevertheless, we expect apoB100 to be mainly present in circulating lipoproteins and possibly in the choriocapillaris.

**FIGURE 4.** Assessment of autofluorescence in apoB100,LDLR–/– mice with SLO. Autofluorescent native images were taken at 488 nm from 14-month-old apoB100,LDLR–/– (a, b at higher magnification) and control mice (c). Perivascular cells were observed at 488 nm in all apoB100,LDLR–/– mice (d). The vascular structure was analyzed in 14-month-old apoB100,LDLR–/– mice in the 514-nm surface image (e) and after fluorescein dye injection at 488 nm (f).
as has been reported. Assuming that LDL can reach RPE and LDLR is present in RPE, the lack of LDLR in apoB100,LDLR / / mice may abrogate the transfer of cholesterol from circulating LDL into RPE cells, leading to cholesterol deposition. Apart from the interaction of LDLR and apoB100 as a pathway for the transfer of lipid and nutrients to the outer retina, other mechanisms may be suggested, as indicated by the presence of CD36 in the RPE. This pathway may be an alternative route for lipid and nutrients to cross the RPE barrier and target the outer retina, in addition to the lack of LDLR, and may explain the non-null ERG response in apoB100,LDLR / / mice, as well as why DHA was not reduced in the neural retina of apoB100,LDLR / / mice, since one would expect low DHA levels if lipid transfer from the RPE to the neural retina was impaired.

Cholesteryl ester deposition at the basement of the RPE (Fig. 5) may contribute to fundus autofluorescence under laser excitation at 488 nm (Fig. 4). Indeed although lipofuscin remains the dominant autofluorescent entity, other lipid deposits may also be detectable in vivo by SLO, in particular in conjunction with OS degradation. Further work is needed to investigate the nature of these deposits, especially the presence of lipofuscin in apoB100,LDLR / / mice.

The autofluorescent perivascular cells (Fig. 4) may be remnants of vitreous vessel walls that have vanished during the course of ocular development. Alternatively, based on their location in the vicinity of inner retinal vessels, these may be lipid-rich smooth muscles cells or pericytes and would correspond to foam cells observed in major aortic vessels during atherosclerosis. Such an explanation remains probable and
accounts for the fact that that we, like others,15 observed fatty streaks in thoracic aorta of these animals (data not shown) and that apoB100,LDLR−/− is recognized as a relevant mouse model for atherosclerosis.15 Accumulation of lipids under the RPE and within BrM is one of the features of aging1,2 and AMD.3,44 AMD is the leading cause of blindness in the elderly in the United States and Western countries.35,40 Various factors have been associated with AMD, including genetics, environmental factors, aging, cardiovascular disease, and risk factors for cardiovascular disease.14,46–49 The mechanisms underlying this phenomenon are being actively investigated, but remain partially resolved. Meanwhile, the consequences for the retinal cycle are poorly described. Using an animal model characterized by systemic lipid dysfuntion triggered by exclusive apoB100 expression and LDLR invalidation, we were able to describe cholesteryl ester deposition in the retina, together with a reduced ERG response. The present apoB100,LDLR−/− and the transgenic apoE4 mouse90 are the only models with neutral lipid deposits at the basement of the RPE that can potentially be useful in studying the mechanisms of lipid deposition that occurs universally in the aging human retina.

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References

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Animal Model for Lipid Metabolism Dysfunction

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