Subretinal Pigment Epithelial Deposition of Drusen Components Including Hydroxyapatite in a Primary Cell Culture Model

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PURPOSE. Extracellular deposits containing hydroxyapatite, lipids, proteins, and trace metals that form between the basal lamina of the RPE and the inner collagenous layer of Bruch's membrane are hallmarks of early AMD. We examined whether cultured RPE cells could produce extracellular deposits containing all of these molecular components.

METHODS. Retinal pigment epithelium cells isolated from freshly enucleated porcine eyes were cultured on Transwell membranes for up to 6 months. Deposit composition and structure were characterized using light, fluorescence, and electron microscopy; synchrotron x-ray diffraction and x-ray fluorescence; secondary ion mass spectrometry; and immunohistochemistry.

RESULTS. Apparently functional primary RPE cells, when cultured on 10-μm-thick inserts with 0.4-μm-diameter pores, can produce sub-RPE deposits that contain hydroxyapatite, lipids, proteins, and trace elements, without outer segment supplementation, by 12 weeks.

CONCLUSIONS. The data suggest that sub-RPE deposition formation is initiated, and probably regulated, by the RPE, as well as the loss of permeability of the Bruch's membrane and choriocapillaris complex associated with age and early AMD. This cell culture model of early AMD lesions provides a novel system for testing new therapeutic interventions against sub-RPE deposition formation, an event occurring well in advance of the onset of vision loss.

Keywords: hydroxyapatite, lipids, x-ray fluorescence, secondary ion mass spectrometry, x-ray diffraction

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ge-related macular degeneration (AMD), the third largest cause of vision loss globally,1 is expected to affect 288 million individuals worldwide by 2040.2 The end stages of AMD are geographic atrophy and neovascular AMD. At present, there is no treatment for geographic atrophy.3 End-stage neovascular AMD can be treated with regular injections of VEGF inhibitors,4 a strategy to prevent progression to late AMD is to modulate deposit formation and clearance. A better understanding of the molecular events leading to the retention of drusen components is essential for the success of this strategy. A model system that recapitulates as many aspects of drusen and BLinD formation as possible would accelerate progress toward these goals.

Sources of the molecular components of drusen and BLinD are not yet established, although candidates exist. An early hypothesis suggests that these lesions are formed from byproducts of photoreceptor outer segment phagocytosis by the RPE.7 However, it was later postulated that a lipid-recycling pathway, driven by diet, of RPE-secreted lipoproteins is responsible.6 Johnson et al.9 showed that highly differentiated and well-polarized primary human foetal RPE cells (hRPE) secreted several deposit-associated proteins without exposure to outer segments or choroidal blood flow. This supports the
idea that some components of deposits originate from the RPE and/or components derived from the culture medium. Proteomic analysis of isolated human drusen also implicated circulating plasma as a protein source.10

Regardless of source(s), it is important to understand what triggers the retention of lipids and proteins in the sub-RPE space. One potential mechanism is binding of RPE-produced lipids and lipoproteins to extracellular matrix elements including proteoglycans in BrM, a mechanism similar to plasma lipoproteins binding to arterial intima in atherosclerosis.11 A second mechanism is retention of proteins via formation of large oligomers,12 a process that is accelerated in the presence of trace metals like zinc.13,14 A third mechanism is binding of proteins to the hydroxyapatite (HAP) spherules recently identified in the sub-RPE space and in drusen of humans.15 Any of these or a combination may be operative.

Published cell culture models for sub-RPE deposit forma-
tion16,17 each identified some, but not all, of the major molecular constituents of human sub-RPE deposits, lipids,18,19 hydroxyapatite,15 trace elements,13,14 and proteins,10,20 in a single model system. A molecularly and physically faithful model system should recapitulate these molecular constituents and form both diffuse and focal deposits. Here, we present evidence that primary RPE cells under standard culturing conditions can create such deposits. Thus, key physical and molecular signatures of AMD lesions, verified by histopathology and high-resolution molecular imaging, can be recreated in vitro by primary RPE cells in a standard medium and without supplementation by either retinal extracts or photoreceptor outer segments.

**Materials and Methods**

**Primary RPE Cell Culture**

Freshly enucleated porcine eyes were recovered from animals following unrelated physiology experiments, transported to the laboratory in ice-cold standard saline, and processed as described.21 In brief, the eye was opened with a cut at the equator that avoided the vitreous body. With one pair of forceps, the anterior half of the eye, including the vitreous body, was lifted away from the posterior half of the eye. The posterior portion contained the retina, RPE/choroid, and sclera. The retina was gently removed with forceps. The RPE cells were released from the eyecups by treatment with trypsin 0.25% and EDTA 0.02% (Gibco, Grand Island, NY, USA). To avoid contamination with other cells, we performed a density gradient centrifugation through a cushion composed of Percoll 40% (Pharmacia Biotech, Piscataway, NJ, USA) with 0.01 mol/L Na2PO4 and 0.15 mol/L NaCl, pH 7.4, at room temperature. After centrifugation, pigmented cells were recovered in the pellet, whereas unwanted cells remained near the top of the cushion. Primary RPE isolated from two porcine eyes were used to establish six Transwell cultures (24-mm polyester membrane, 0.4-μm pore size; Costar, Corning, NY, USA) without laminin or collagen coating. Cells were cultured in Miller medium22 containing 1% fetal calf serum for up to 6 months, per the results. Medium was changed every 3 to 4 days. When applicable, primary cell cultures were subcultured every 2 weeks at a ratio of 1:2, meaning that one dish was divided into two identical dishes and then maintained under the same conditions.

**Human Fetal RPE Cell Culture**

Primary human fetal RPE cells (hRPE; Sciencell, Carlsbad, CA, USA) were propagated in a T-75 flask (Corning; Thermo Fisher Scientific, Waltham, MA, USA) with Epithelial Cell Medium (EpiCM) (Sciencell), completed with 2% fetal bovine serum (Sciencell), 1:100 mL/mL epithelial cell growth supplement (Sciencell), and 107 U/L penicillin/10 g/L streptomycin (Sciencell). For our experiments, cells at passage 2 were cultured either on 10-μm thick Transwell inserts with a pore diameter of 0.4 μm (Corning; Thermo Fisher Scientific) coated with Geltrax extracellular matrix (Thermo Fisher Scientific) or on 100-μm thick Transwell inserts with a pore diameter of 0.45 μm (Millipore; Thermo Fisher Scientific) coated with laminin (Sigma-Aldrich Corp., St. Louis, MO, USA), in MEM-x modifications as previously described.22

**Immunohistochemistry of Porcine Cells**

Immunofluorescence was performed on flat mount and cross sections of porcine RPE cells using methods for cells on coverslips.21 Cross sections shown in Figure 1 were 10 μm thick and were generated using a cryostat (Leica CM 3050 S; Leica, Bannockburn, IL, USA). Primary antibodies included cytokeratin 18,23 zonula occludens,24 apically localized Na+-K+ ATPase,25 cellular retinaldehyde binding protein,26 and microsomal triglyceride transfer protein.27 For apolipoprotein E (ApoE) staining, cultures were fixed in 4% paraformaldehyde with 1% glutaraldehyde, cryopreserved in 30% sucrose overnight, and then embedded in optimal cutting temperature compound (Agar Scientific, Stansted, Essex, UK). Cryosections, 10 μm thick, were generated using a cryostat (Leica CM 3050 S; Leica) and collected on glass slides. Cryosections were blocked with a 1:20 dilution of donkey serum in PBT (D9663; Sigma-Aldrich Corp.) for 1 hour at room temperature. Subsequently, sections were incubated with primary antibody followed by secondary antibody for 1 hour each. The primary antibody used was goat anti-ApoE polyclonal (1:500 dilution; #AB947; Millipore, Billerica, MA, USA) and the secondary antibody was an Alexa Fluor donkey anti-goat secondary antibody (1:200 dilution; excitation wavelength, 488 nm; A11055; Molecular Probes). To visualize nuclei, Hoechst nuclear stain (1:5000 dilution; excitation wavelength, 358 nm; H21492; Molecular Probes) was used. Samples were incubated for 20 minutes at room temperature and in the dark. Images were generated using a Zeiss LSM700 confocal microscope (Oberkochen, Baden-Württemberg, Germany) through a 63×/1.2 NA Zeiss Neofluar objective.

**Immunohistochemistry of Human Fetal RPE Cells**

Immunohistochemistry was performed according to the procedure described for porcine cells, except that human fetal RPE cells were cultured for 8 weeks and fixed in 4% paraformaldehyde before embedding and sectioning. Primary antibodies used in these experiments were against ApoE (1:200 dilution; #AB947; Millipore) and/or ZO1 (1:400 dilution; 1:400 dilution, 610966; BD Biosciences, Franklin Lakes, NJ, USA). Alexa Fluor 546-561 and 488 IgG (H+L) (1:200; 1:1056 and 21202; Molecular Probes) were used as secondary antibodies to detect anti-ApoE and anti-ZO1, respectively.

**Synchrotron Microfocus X-ray Fluorescence and Microfocus X-ray Diffraction Analysis**

Synchrotron microfocus x-ray diffraction (μXRD) and microfocus x-ray fluorescence (μXRF) analysis of cell cultures was collected at the 15ID-E beamline at the Advanced Photon Source (APS), Argonne National Laboratory. Analytical methods and instrument configuration follow those described in Lanzirotti et al.28 Porcine cell cultures were flat mounted on Kapton film for analysis. An incident beam energy of 17.48 keV
was used, focused to a spot size of ~1.3 \text{\mu m} (H \times V) with a photon flux of ~10^{11} \text{photons/s} within this focused spot. X-ray fluorescence from the sample was measured using a four-element Vortex ME4 (Hitachi, Inc., Northridge, CA, USA) silicon drift diode detector coupled to an Xspress 3 digital x-ray multichannel analyzer system (Quantum Detectors, Harwell, Oxford, UK). X-ray diffraction from the sample was measured using a Perkin Elmer XRD1621 digital flat panel detector (Santa Clara, CA, USA) placed in transmission geometry.

Note, experiments should control for the artifactual appearance of XRD signatures from the buffer systems used (Supplementary Fig. S1).

**Identification of HAP, Protein, and Lipid by Time of Flight–Secondary Ion Mass Spectrometry Imaging**

Secondary ion mass spectrometry (SIMS) and secondary ion mapping was performed using a TOFSIMS5-Qta\textsuperscript{100} LEBS mass spectrometer (ION-TOF, Münster, Germany). Submicrometer resolution was used for secondary ion mapping over m/z 0-880. The system is comprised of a bismuth primary ion beam, operating at 25 kV and tuned to use the Bi\textsuperscript{+} cluster for greater secondary ion yield, and a low energy electron flood gun for charge compensation. Ionic species sputtered from the surface under the bismuth bombardment are steered into a reflectron time-of-flight mass analyzer. Before mass spectrometry was performed, an increased ion beam current was used to remove potential contamination from the upper surface of the sample. Identified peaks strongly localized to deposits were mapped on single ion maps.

**Hydroxyapatite Fluorescent Labeling**

Fixed and de-waxed paraffin embedded sections, 20 \text{\mu m} thick, with crystalline deposits were stained with 20 \text{\mu M} bone-tag 680RD (926-09374; Li-Cor) or 20 \text{\mu M} OsteoSense (NE-V10020EX; Perkin Elmer) in aqueous buffer for 20 minutes at room temperature. Samples were imaged using a Zeiss LSM700 confocal microscope through a 63\times/1.2 NA Zeiss Neofluar objective with an excitation wavelength of 620 nm and emission wavelength of 680 nm.

**Transmission Electron Microscopy**

Culture dishes were preserved in cold 1% glutaraldehyde, 2.5% paraformaldehyde, in 0.1 M phosphate buffer and photodocumented using a dissection microscope with a dark-field illumination base. Transwell inserts with cells attached were cut into 2 \times 2 squares, photodocumented for orientation, postfixed by the osmium tannic acid paraphenylenediamine method to accentuate neutral lipid, sectioned at silver-gold thickness, and viewed with a 1200 EXII electron microscope (JEOL USA, Peabody, MA, USA) and an AMTXR-40 camera (Advanced Microscopy Techniques, Danvers, MA, USA).

**Scanning Electron Microscopy**

Fixed cell cultures stored in PBS were dehydrated in a graded ethanol series (20%, 50%, 70%, 90%, and 100%) before critical point drying in hexamethyldisilazane (Sigma-Aldrich Corp.). Samples were mounted using carbon adhesive tabs on aluminium stubs (Agar Scientific). Specimens were sputter coated with gold/palladium and imaged using a Philips XL30.
FEG-SEM (Hillsboro, OR, USA) with a spot size of 3 and an accelerating voltage of 5 kV.

Microcomputed Tomography

Samples were dehydrated as described for scanning electron microscopy. Dehydrated samples were mounted on to a rotary stage using an adhesive. Microcomputed tomography (micro-CT) was performed within a Philips XL30 FEG-SEM using the SkyScan \textsuperscript{\textregistered} CT system (Bruker System, Billerica, MA, USA). Samples were scanned every 1.0\textdegree through 180.0\textdegree. Resolution was 1 \textmu m per pixel. Three-dimensional reconstruction of scanned slices was achieved by volume rendering using the CTVox reconstruction software (Bruker System).

Human Eye Pathology

The Institutional Review Board at the University of Alabama at Birmingham approved the use of human tissue. Short postmortem (<6 hour) donor eyes used in prior publications were reviewed to identify deposits containing lipids, refractile material, and von Kossa–positive material. These include 82 AMD maculas (53 late AMD and 29 early AMD) processed for high-resolution submicrometer sections for the Project MACULA website of AMD histopathology\textsuperscript{29,30} (http://projectmacula.cis.uab.edu/); geographic atrophy eyes (n = 10) stained with the von Kossa method\textsuperscript{31}; and early and advanced AMD eyes (n = 12) stained for lipids and with the von Kossa method.\textsuperscript{32,33} Selected areas were imaged using 60\times oil immersion objectives (numerical aperture, 1.40). Images were montaged, composited, and adjusted for exposure, contrast, and sharpness (Photoshop CS6; Adobe, San Jose, CA, USA).

Study Approval

Use of porcine eyes was approved by the Institutional Animal Use and Care Committee at the University of Alabama at Birmingham.

RESULTS

Primary Cell Cultures Display Characteristics of Well-Differentiated Cells

At 1 day after isolation, clumps of heavily pigmented cells were attached and well spread on porous polyester Transwell membranes (Costar). Seven days after isolation, rapidly proliferating cells formed larger clumps and maintained cell–cell contacts and epithelioid morphology. By 14 days and thereafter, still-polygonal cells formed a confluent monolayer with high (>150 MO) trans-epithelial resistance, a comprehensive measure of RPE polarity, confluence, and integrity of cell–cell contacts. Based on morphology, adhesion, and proliferation, cells were characterized as primary RPE cells. Whether examined en face or as vertical cryo-
sections (Fig. 1, left and middle columns), confluent monolayers showed strong immunoreactivity for markers of intermediate filaments, junctional complexes, polarized ion transporters, retinoid processing, and lipoprotein assembly, which are features that typify RPE in situ.

These primary cultures were associated with intermittent patches of lipid and phosphate deposition revealed by classic histochemical stains (Fig. 1, right column). Cells at 6.0 weeks were hexagonal especially in areas with high melanosome content (Fig. 1, Melanin). At this time, staining with oil red O was prominent near the pigmented cells (Fig. 1, Oil red O). von Kossa staining produced a patchy light brown reaction product suggesting mineralization was also present (Fig. 1, von Kossa).

Examination of Sub-RPE Deposits With Light, Fluorescent, and Electron Microscopy

Cells were well differentiated, as evidenced by numerous apical processes (Figs. 2A, arrow, 2B). A layer of toluidine blue-stained material between the RPE cells and the cell culture insert was observed (Fig. 2A). This material was electron dense (Fig. 2B) and unaccompanied by thickening of the basal lamina. Additionally, intense oil red O staining confirmed the presence of neutral lipids within the deposits and within the pores of the culture dish insert (Fig. 2C). Staining of pores was not detectable in inserts of dishes lacking cells. Electron microscopy also revealed the presence of cellular processes and material of varying electron density within pores (Fig. 3).

Fluorescent dye staining for HAP indicated varying degrees of mineralization below the RPE (Figs. 2D, 2E). In all samples, deposits were confined to the sub-RPE compartment without equivalent deposits on the apical surface. Deposits were continuous on the spatial scale of several hundred cells but not across the entire dish and, as such, are referred to as diffuse.

Analysis of Sub-RPE Deposits With Electron Microscopy and Micro-CT

Next, we examined deposits at higher resolution using transmission electron microscopy (Figs. 4A–D) and scanning electron microscopy (Figs. 4E–H). Additionally, micro-CT was used for detection of mineralization (Figs. 4I–L). By following cultures over time, showed it was determined that passaged cells did not develop deposits under these experimental conditions, and thus these cells are designated deposit-incapable. Cell cultures were examined at three time points in culture and in two differentiation states: 6.0 weeks (deposit-incapable third passage cells), 12.0 weeks (primary cells with deposits), and 26.5 weeks (primary cells with abundant focal and diffuse deposits). Deposit-incapable passaged cells cultured for 6.0 weeks lacked apical processes, tight junctions, melanosomes, deposits, and signal for mineralization (Figs. 4A, 4E, 4I). Deposit-capable primary cells at 6.0 weeks exhibited sparse, short, and stubby apical processes and formed apically located...
tight junctions and variable melanosome content. The cells also formed deposits that were not readily visible by transmission electron microscopy (Fig. 4B) but were clearly detected by micro-CT (Fig. 4J). Deposit-capable primary cells at 12.0 weeks exhibited little or no mineralization. N, nucleus; Ins, culture dish insert. (B, F, J) Deposit-capable primary cells at 6 weeks produced a thin electron-dense sub-RPE deposit that was continuous with material in pores (arrow) and exhibited some mineralization. Due to the sectioning plane, the pore does not cross the Transwell. (C, G, K) At 12 weeks, deposit-capable cells developed a continuous layer of deposit with increased mineralization. (D, H, L) At 26.5 weeks, focal dome-shaped deposits were also present, with diffuse deposit, exhibited intense mineralization (H, I; arrowheads). Diffuse and focal deposits had solid cores (yellow asterisks) with feathery surfaces (red arrowhead) (C, D). Cracks between cells (G, H) are due to brittleness conferred by mineralization. Scale bars denote the following: A–D, 2 μm; E–L, 100 μm.

**Elemental and Mineral Characterization of Deposits Using µXRF and µXRD**

We analyzed the elemental composition of cell culture deposits using µXRF (Fig. 6). The deposits produced by the cultured RPE cells were enriched in Ca (Figs. 6B, 6C), Fe (Fig. 6D), and Zn (Fig. 6E). Furthermore, the mineral content of deposits was determined through implementation of µXRD. The summed µXRD patterns (Fig. 6H) and corresponding radial intensity profiles (Fig. 6G, blue line) clearly identified the presence of HAP compared with a reference HAP 2h profile (Fig. 6G, red bars). Mapping mode µXRD of the integrated intensity peaks representing the HAP 002 and 211 crystallographic reflections (Figs. 6I, 6J) showed that HAP was present throughout all examined focal deposits. Mapped HAP reflections were also observed between focal deposits, suggesting the presence of diffuse deposits within the examined area. These data provide evidence that primary RPE cells are capable of developing HAP-containing sub-RPE deposits that also contain Fe and Zn.

**Characterization of Deposits Using SIMS**

To determine whether deposits in culture also contain other constituents of human sub-RPE deposits, we analyzed sections of cultured cells containing visible focal deposits using SIMS. Typical ionic signatures for HAP (Supplementary Fig. S2A) and proteins (Supplementary Figs. S2B–S2E; represented by amino acids) were readily identifiable within nodules. Secondary ion
Drusen in a Dish

**DISCUSSION**

Herein, we demonstrated that well-differentiated primary porcine RPE cells in culture produce focal and diffuse sub-RPE deposits that contain HAP, lipids, proteins, and trace metals, as in deposits in human AMD macula. Deposits have been an elusive aspect of AMD pathology in model systems to date. To our knowledge, this is the first time that these four major molecular constituents have been replicated and confirmed in a single cell culture system, albeit with physical forms resembling, but not yet identical to, sub-RPE deposits in vivo. These achievements are significant for the molecular basis of AMD’s specific lesions, RPE cell biology, and the prospects of a high-fidelity model system suitable for basic and translational AMD research.

Our experiments used 10-μm-thick polyester membrane inserts that imposed a physical barrier against movement of RPE secreted material or material present in the basal compartment of the culture medium. Our observations can be compared and contrasted with those of Fernandez-Godino et al., who reported that primary mouse RPE cells after 2 weeks in culture released material into 0.4-μm-diameter pores connecting apical and basal chambers, with cellular processes ramifying on the opposite side of the insert. Although we occasionally observed cellular processes in pores (Fig. 3), we observed extracellular deposits primarily between the RPE cells and the insert. Amin et al. showed extracellular material between ARPE-19 cells and a smooth surface without pores or a basal compartment after 11 weeks. Using fetal human RPE cells on 100-μm-thick porous supports (PIHA 01250; Millipore), Johnson et al. demonstrated particulate deposition of ApoE-immunoreactive secreted material exclusively within the insert and not on the insert surface. We also achieved particulate ApoE-immunoreactive deposition by human fetal RPE when using an identical insert. Importantly, when using a 10-μm-thick polyester membrane, we showed that these same cells produced extensive deposits on the insert surface (Fig. 8D).

Differences among cell culture systems are important to consider when extrapolating to the in vivo situation. Our data highlight the role of physical entrapment of material by aged BrM. In the eye, the “insert” to which the RPE basal lamina attaches is the inner collagenous layer of BrM. In aging macula, inner and outer collagenous layers of BrM thicken and the elastic lamina calcifies. Once lipids and calcium are deposited, BrM could act as a physical barrier that retains proteins and lipids between the RPE basal lamina and inner collagenous layer. In this environment, entrapped lipids, calcium, and phosphate could promote HAP spheroide formation, to which monomeric and/or oligomerized proteins could bind. It will be interesting to determine whether manipulation of the insert surface will promote deposit formation. Although our data indicates that a barrier is essential for formation of deposits, our studies do not address the relative contributions of aging BrM and choriocapillaris to this barrier. Cocultures of RPE and choriocapillary endothelium may thus prove informative.

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binding lipid (including esterified cholesterol) in soft drusen. In contrast, deposits reminiscent of subretinal drusenoid deposits (between apical RPE and photoreceptors lacking oil red binding lipid and refractile bodies; Fig. 9A), BLamD (Fig. 9B), and end-stage drusen (aggregated refractile, von Kossa-positive nodules without histochemically detectable lipid; Figs. 9E, 9F) were absent from our culture system. These differences (Supplementary Table S1) underscore that further optimization is required to recapitulate all aspects of in vivo deposit formation. Fortunately, factors such as extracellular pH, concentration of extracellular calcium, phosphate, and trace metals and protein environments, and external factors such as supplementation can be easily tested and optimized in a cell culture system.
There are two theories of drusen biogenesis: transformation (direct conversion of RPE cells to druse material) and deposition (material released by an otherwise intact RPE cell layer). Our data support the latter theory, with the addition that material of RPE origin accumulates due to an outflow barrier in subjacent BrM. The initial motivation for these experiments was the hypothesis that RPE secretes ApoB and E-containing lipoprotein particles in a constitutive and physiologic recycling system to dispose of excess lipids delivered by diet. Cell autonomous secretion of lipid in our system, along with previously reported secretion of ApoE-immunoreactive lipoprotein-like particles without outer segment supplementation, supports an overall physiologic and diet-driven model. This model is also supported by recent clinical imaging studies showing survival of the RPE layer over soft drusen for multiple years. Over this period, drusen increase in volume and then collapse after the RPE layer disintegrates.

In the porcine RPE system, early passage inhibited the ability of cells to form deposits, likely due to irreversible loss of phenotype via transdifferentiation. This finding has implications not only for RPE model systems such as immortalized cell lines like ARPE-19, but also for replacement cells being considered for therapeutic transplantation. Replacement cells might create deposits in the eye with deleterious consequences like those of native deposits. Conversely, transplanted cells that cannot produce deposits on an appropriate substrate may lack key cellular functions.

We demonstrated continuous sub-RPE deposits in the absence of many factors thought to be important in druse initiation. These include photoreceptor outer segments, human-specific serum components, non-RPE cells such as macrophages, RPE lipofuscin, BLamD, stress beyond that associated with isolation, exogenously applied injury, aging, and light exposure. Other factors not required are the gene for cholesterol ester transfer protein, an AMD risk factor that

Figure 7. Distribution of the molecular ions identified by SIMS. An optical image of the focal deposit analyzed by SIMS analysis (A). Molecular ion mapping of calcium (Ca\(^{2+}\)) (B) and calcium phosphate (Ca\(_2\)PO\(_4\)\(^{-}\)) (C) confirmed the presence of HAP within sub-RPE deposits. Lipid content of deposits was confirmed by molecular ion mapping of the phosphatidylcholine (PC) head group fragment ion (D). Mapping of amino acid signatures including glycine (E), alanine (F), proline (G), and leucine (H) confirmed the presence of protein within deposits. Thus, HAP amino acids, and PC are all present within deposits. It should be noted that the culture insert is absent in the figure, during processing the cells and the sub-RPE deposits detached from the Transwell insert surface. Scale bars denote 40 μm.

Figure 8. Accumulation of ApoE in sub-RPE deposits in primary porcine and human fetal RPE cell culture models. Immunohistochemistry was performed on porcine RPE cells cultured for 26.5 weeks (A–C) or on human fetal RPE cells cultured for 6 weeks (D, E). Transwell membranes were either polyester (A–D) or composed of mixed cellulose (E). (A) Punctate (arrowbeads) and diffuse ApoE immunofluorescence was present within focal deposits, as well as on the surface of the polyester insert when cells were cultured for 26.5 weeks. (B) Apolipoprotein E-immunofluorescent deposits accumulated along the membrane in a nonuniform manner (arrow). Note the intense labelling within the RPE cell bodies and less intense labeling within pores of the Transwell membranes on A and B. (C) Punctate ApoE labeling (arrows) is found within or under HAP sub-RPE deposits (magenta). (D) Human fetal primary RPE cells grown for 8 weeks on polyester membrane inserts show ApoE (green) accumulation on the surface of the membrane, similar to that for porcine cells. Unlike porcine cells, human RPE cell bodies stain lightly for ApoE. Red immunofluorescence corresponds to ZO1 staining associated with junctions, indicating that the cells are well differentiated. (E) Human cells grown on membranes of mixed cellulose esters showed the dispersed accumulation of ApoE (green) particles, identical to those published by Johnson et al. in the membrane cavities. Red arrows between C and E indicate the surface of the membrane inserts. Scale bars denote 10 μm in A–E.
is absent in pigs,59 or any specific genotype. Without excluding direct deposition of culture medium components into the sub-RPE space, we can hypothesize that the minimum requirements for deposits are functional RPE cells that can secrete lipoproteins and molecules required for HAP formation, plus a retentive barrier.

The multiple molecular commonalities between in vitro deposits and human drusen provide impressive biological plausibility of this cell culture system. Questions to be addressed in future studies are generalizability to RPE from other species, time course of deposition, impact of outer segment supplementation, and a comprehensive molecular comparison of in vitro deposits versus human AMD lesions. Such data would support use of this model system for testing the regulation and therapeutic abatement of sub-RPE deposits.

This is important because deposits provide a proinflammatory proangiogenic nidus and cleavage plane for neovascularization.60 Reduction of drusen volume61 through targeted therapies62 has the potential to mitigate the treatment burden of VEGF suppression for neovascular AMD. Our culture system may be useful in achieving this goal.

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CAC guarantees the integrity of the experiments, results, and analysis in this study and had full access to all of the data generated.

CAC and CG hold intellectual property related to this manuscript. The remaining authors declare that no conflict of interest exists.

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