ABUNDANCE AND MULTIMODAL VISIBILITY OF SOFT DRUSEN IN EARLY AGE-RELATED MACULAR DEGENERATION

A Clinicopathologic Correlation

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Purpose: To determine the abundance and multimodal visibility of drusen and basal linear deposit (BLinD) in early age-related macular degeneration.

Methods: A 69-year-old white man was imaged by color fundus photography and red free photography, fundus autofluorescence, and optical coherence tomography. From en face images, we determined the drusen field, drusen area, and equivalent diameters of individual drusen. From high-resolution light-microscopic histology (6 months after the last clinic visit), we determined the area of drusen, BLinD, and pre-BLinD in a subretinal pigment epithelium-basal lamina lipid field.

Results: In right and left eyes, respectively, BLinD covered 40% and 46% of the lipid field, versus 21% and 14% covered by drusen. The lipid field was covered 60% to 61% by Drusen + BLinD and 65% to 72% by BLinD + pre-BLinD. In the left eye, the drusen area on color fundus photography (0.18 mm²) and red free (0.28 mm²) was smaller than the drusen area on histology (1.16 mm²). Among drusen confirmed by optical coherence tomography, 55.1% and 56.6% were observed on red free and fundus autofluorescence, respectively.

Conclusion: Basal linear deposit covered 1.9 and 3.4-fold more fundus area than soft drusen, silently increasing progression risk. Improved visualization of BLinD and readouts of the retinal pigment epithelium health over lipid will assist population surveillance, early detection, and trial outcome measures.

Drusen are the largest intraocular risk factor for progression to advanced age-related macular degeneration (AMD). Longitudinal population-based studies using color fundus photography (CFP) demonstrate steady accumulation and growth of drusen from midlife, concentrated in the central macula.¹ Soft drusen and basal linear deposit (BLinD) are lump-and-layer versions of the same lipid-rich extracellular material located between the basal lamina of the retinal pigment epithelium (RPE-BL) and the inner collagenous layer of Bruch membrane (BrM) (Figure 1).² A well-supported model of drusen formation posits that RPE secretes lipoproteins constitutively to offload unneeded lipids from dietary uptake and photoreceptor outer segments, and this process is impeded by poor transfer across aged BrM and choriocapillaris.³ Soft drusen material is a direct precursor to AMD end stages Type 1 neovascularization (Figure 1) and atrophy.

Accurate quantification of soft drusen for progression risk estimates in populations and early AMD trial outcomes is doubly challenged. First, BLinD may confer similar risk as soft drusen (Figure 1) yet is not yet routinely visible either by clinical imaging or histology. Second, multimodal imaging including optical coherence tomography (OCT) has established the
presence of the subretinal drusenoid deposit (also called reticular pseudodrusen), which may have been counted as drusen in epidemiology studies.4

Clinicopathologic correlation of eyes imaged during life provided essential insight into sub–RPE-BL lipid.2

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By analyzing through high-resolution histology fellow eyes with early AMD and in vivo multimodal OCT-based imaging, we quantified the proportion of the macula containing drusen and BLinD. We also assessed pre-BLinD (Figure 1), a layer of lipoprotein particles in the sub–RPE-BL space that forms after the 3-layer substance of BrM fills up throughout adulthood. Thus, for the first time in this one case, all forms of high-risk lipid could be assessed. Not only is invisible BLinD more abundant than soft drusen but also not all drusen are visible en face. Data highlight the need for outcome assessments incorporating all lipid-rich deposits.

Methods

A 69-year-old HIV-positive white man presenting with early AMD underwent color fundus and red free (RF) photography (Topcon TRC 50DX; Topcon Medical Systems, Oakland, NJ), spectral domain OCT (30° × 20°, 19 scans, 240 μm spacing, Automated Real-time Averaging 10–11; Heidelberg Spectralis HRA + OCT, Heidelberg Engineering, Heidelberg, Germany), and fundus autofluorescence (FAF). Color fundus photography and RF were acquired 14 months and OCT and AF 6 months before death. Color fundus photography, RF, and FAF were registered to the near-infrared reflectance image by vascular landmarks and then linked to OCT. Eyes preserved 9 hours after death were postfixed in osmium tannic acid paraaffinembedded in epoxy resin, sectioned at 0.8 μm, and stained with toluidine blue. From

![Image](image-url)
glass slides of the entire left macula and the inferior half of the right macula, one section per slide was scanned with 20X and 60X oil immersion objectives (Olympus VSI 120, CellSens; Olympus, Center Valley, PA).

Measurements defined in Supplemental Digital Content (see Table 1, http://links.lww.com/IAE/B270), used ImageJ (https://imagej.nih.gov/ij/download.html) except where noted. We measured histologic cross-sectional lengths of BrM covered by different categories of sub–RPE-BL lipid (Figure 2, A–E). Pre-BLinD is thin, flat, gray, and finely granular. Basal linear deposit is thin, undulating, with loose osmophilic material like soft drusen. Pre-BLinD and BLinD may be continuous with yet distinct from drusen, which are dome shaped. To determine the BrM area covered by the lipid, we multiplied cross-sectional lengths along BrM by intersection distances (Cavaliere’s principle). The full extent of the histologic lipid was considered the lipid field.

In clinical CFP and RF images, we measured the drusen field, a closed contour bounding visible drusen (Figure 2, A and B and D, E). The drusen area was the sum of all individual drusen (see Figure 1 A1–D1, Supplemental Digital Content, http://links.lww.com/IAE/B269). Equivalent diameters of individual drusen were calculated assuming each drusen had a circular base. Lengths of drusen along BrM on OCT B-scans were measured using the distance measurement tool of the Spectralis. Equivalent diameters are slightly larger than cross-sectional lengths and are reported separately. We evaluated whether drusen visible on OCT were also visible in en face imaging. To assess whether ex vivo measurements were impacted by tissue volume changes during processing, we determined that the distances between the fovea and optic nerve head edge were similar in histology and clinical imaging.

The clinical imaging study was approved by the institutional review boards of the University of California, San Francisco and the University of California, Berkeley. The Foundation Fighting Blindness eye donor program provided the tissue specimens. The histopathology study was approved by the University of Alabama at Birmingham, complied with the Health Insurance Portability and Accountability Act of 1996, and adhered to the tenets of the Declaration of Helsinki.

Results

Table 1 shows histologic abundance of different categories of the sub–RPE-BL lipid. Figure 2F shows

Fig. 2. Distribution of categories of the sub–RPE-BL lipid in early AMD. A. A panoramic view of a section showing categories of the sub–RPE-BL lipid. Black dashed frames are magnified in panels B–E. B. No sub–RPE-BL lipid. C. Pre-BLinD is a flat layer of finely granular material in gray (green arrowheads). D. Basal linear deposit is an undulating layer of the same extracellular material as in soft drusen (yellow arrowheads), usually continuous with pre-BLinD. E. Soft drusen are lump version of the same extracellular material, usually continuous with BLinD. F. Sub–RPE-BL lipid distribution in fellow eyes. AMD, age-related macular degeneration; BL, basal lamina; BLinD, basal linear deposit; BLamD, basal laminar deposit; Ch, choroid; ChC, choriocapillaris; d, drusen; OS, outer segment. All lipid, drusen + BLinD + pre-BLinD. Scale bar in B applies to B–E.
the percentage of BrM covered by these lipid forms. In the right and left eyes, respectively, Drusen + BLinD, together comprising the high-risk lesion of AMD,\(^6\) accounts for 2.81 mm\(^2\) to 5.08 mm\(^2\) (60–61%) of the lipid field. Clinically invisible BLinD + pre-BLinD account for 2.99 mm\(^2\) to 6.17 mm\(^2\) (65–72%) of the lipid field. Drusen account for only 0.97 mm\(^2\) to 1.16 mm\(^2\) (14–21%). Thus, the area of BLinD was 1.9 to 3.4 times larger than the drusen area.

On CFP and RF clinical imaging, the drusen field was 5.44 mm\(^2\) and 5.61 mm\(^2\), respectively, in the right eye (Figure 3, A and B) and 5.01 mm\(^2\) and 5.35 mm\(^2\) in the left eye (Figure 3, D and E), substantially lower than in histology. Furthermore, the drusen area on CFP and RF were 0.43 mm\(^2\) and 0.55 mm\(^2\), respectively, in the right eye and 0.18 mm\(^2\) and 0.28 mm\(^2\) in the left eye, both smaller than the drusen area on histology. Figure 3, H and I show drusen sizes in four different modalities in the left eye. Drusen detected on CFP and RF are mainly small (31–63 μm) and mainly medium (64–125 μm) on OCT. Interestingly, 19% and 31% of drusen were large (>125 μm) on OCT and histology, respectively, larger than any drusen seen on CFP or RF.

Optical coherence tomography revealed in both eyes fine stripes of hypertransmission, not confined to drusen but prominent at some (Figure 3, C and F). These are attributed to degenerating RPE with fewer shadowing organelles in the light path.\(^7\) Of 127 drusen confirmed by OCT in right and left eyes, 70 (55.1%) were visible within 80 μm of the same locations on RF and 57 (44.9%) could not be found. In the left eye (Figure 3G) were small tiny hypoautofluorescent dots with or without a faint circumferential ring of increased FAF.\(^8\) Among 53 drusen confirmed in OCT B-scans in the left eye, 30 (56.6%) were visible on FAF and 23 (43.4%) could not be found.

### Discussion

For the first time, the full extent of the histologically identifiable sub–RPE-BL lipid in AMD has been

![Fig. 3. Sizes of drusen detected in clinical imaging and in histology. A, B, D, and E. Dashed lines delineate the drusen field in CFP (A and D) and RF (B and E) images. Red free image shows drusen better than CFP. Color fundus photography and RF were acquired 14 months before death and OCT and AF 6 months before death. C and F. OCT B-scan through the fovea shows RPE elevations (yellow; see inset). Stripes of hypertransmission (green), indicating RPE degeneration, are not confined to drusen but are prominent at some. G. AF of the left eye shows multiple tiny hypoautofluorescent dots in temporal macula (green dashed rectangle; magnified in inset). H. Distribution of equivalent diameters of drusen on CFP and RF in the left eye. I. Distribution of cross-sectional lengths of drusen on OCT and histology in the left eye. AF, autofluorescence. L, large drusen; M, medium drusen; S, small drusen.](image-url)
quantified. Basal linear deposit, which is invisible to multimodal clinical imaging, covered 1.9 to 3.4 times more fundus area than soft drusen. Thus, BLinD could substantially but silently increase progression risk. Drusen seen en face by CFP, RF, and FAF appear smaller than seen by OCT, likely because drusen visibility depends on the degree of associated RPE change. FAF signal variation in the drusen field included some definite hypoautofluorescence largely because of lipofuscin loss and rearrangement within dysmorphic RPE.9

Drusen are quantifiable with commercially available OCT. Stabilization or diminution of the drusen volume is proposed as part of a composite outcome measure for clinical trials of therapies targeting early and intermediate AMD stages.10 This attractive concept is challenging to implement because of the dynamism of drusen. Our new data further show that drusen may be only a fraction of an overall lipid field, a relevant consideration if lipid detoxification or removal is the goal.11 One inference from our data is that rather than the directly measure sub–RPE-BL lipid, the impact of that lipid could be assessed with a metric of RPE health. This idea is supported by both the pinstripe hypertransmission seen on OCT and the delicate hypoautofluorescence seen in this case (Figure 3, C and F). In this scenario, a suitable trial endpoint could be stabilization of hypertransmission in the absence of progression.

Our data also suggest that for the future, techniques for visualizing soft drusen material could be improved. Existing, prototype, and experimental imaging technologies with such capabilities12–14 should be prioritized for study and validation. Recently, hypofluorescence in late-phase indocyanine green angiography was proposed as a candidate imaging correlate for BLinD because soft drusen are also hypofluorescent.12 Supporting this idea was an age-related increase in hypofluorescence across the macula.

Study strengths include quantitative, high-resolution, and comprehensive histology of early AMD eyes with in vivo multimodal imaging. Limitations include limited generalizability because of the study of one case, varying time points for imaging modalities, and known imprecision in clinical image registration.15 In conclusion, BLinD covers an area 1.9 to 3.4 times larger than soft drusen and may silently increase AMD progression risk. Not all soft drusen are clinically visible; more are seen with OCT than fundus photography, which may show primarily depigmentation associated with drusen. As new treatments become available, population surveillance and early detection of AMD will become imperative. Progress toward these goals will be aided by improved visualization of BLinD and in the near term, RPE health over lipid.

**Key words:** age-related macular degeneration, autofluorescence, basal linear deposit, clinicopathologic correlation, color fundus photography, drusen, histology, optical coherence tomography.

**References**