Aceruloplasminemia

Retinal Histopathologic Manifestations and Iron-Mediated Melanosome Degradation

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Objective: To examine the retinal histopathologic manifestation of aceruloplasminemia, an autosomal recessive disease caused by mutation of the ferroxidase ceruloplasmin, resulting in tissue iron overload.

Methods: The morphologic features of the human aceruloplasminemic retina were studied with light and electron microscopy. Retinal iron accumulation was assessed with Perls Prussian blue staining, immunohistochemistry, and secondary ion mass spectrometry.

Results: Light and electron microscopic analysis revealed several ocular pathologic findings that resembled age-related macular degeneration, including retinal pigment epithelium (RPE) depigmentation, atrophy and hypertrophy, nodular and diffuse drusen, and lipofuscin and melanolipofuscin granules. Complement deposition was detected in drusen. The RPE cells and neural retina had increased levels of iron. Two major types of RPE cells were observed: melanosome rich and melanosome poor. Melanosome-rich cells had increased levels of iron and melanolipofuscin. The melanolipofuscin granules were observed in large aggregates, where some of the melanosomes were degrading. Melanosome-poor cells lacked melanosomes, melanolipofuscin, and lipofuscin but contained electron-dense aggregates high in iron, phosphorus, and sulfur.

Conclusions: The findings in the aceruloplasminemic retina resemble some of those found in age-related macular degeneration. Also, they suggest that melanosomes in the RPE can be degraded via iron-mediated reactive oxygen species production.

Clinical Relevance: Mechanisms underlying the pathologic mechanisms found in aceruloplasminemia also may be important in age-related macular degeneration.


Aceruloplasminemia is a rare autosomal recessive disease caused by mutation of the ceruloplasmin gene. Aceruloplasminemia is characterized by a triad of diabetes mellitus, retinal degeneration, and neurodegeneration that usually manifests between the ages of 30 and 55 years. The disease is caused by iron accumulation in the pancreas, liver, brain, and retina due to a decrease in iron export.

Ceruloplasmin is a ferroxidase protein that is secreted from cells or is glycosylphosphatidylinositol linked to the outer surface of cell membranes. Ceruloplasmin interacts with ferroportin, the cellular iron exporter, to export ferrous iron from cells. As a ferroxidase, ceruloplasmin converts ferrous to ferric iron, allowing ferric iron to bind to transferrin in the plasma. Lack of ceruloplasmin, even in the presence of ferroportin, markedly impairs iron export from cells. Thus, ferroxidases are necessary components for iron export.

In retinas with age-related macular degeneration (AMD), iron and transferrin levels are elevated, suggesting that iron toxicity may contribute to the pathogenesis of AMD. Furthermore, in mice, combined knockout of ceruloplasmin and mutation of its homologue, hephaestin, causes retinal and brain iron accumulation and degeneration. This retinal degeneration resembles AMD.

Pathologic studies of many aceruloplasminemic tissues have been published; however, no study has examined retinal histopathologic manifestations, to our knowledge. The only published reports focusing on the retina have been clinical case reports, which have shown pigmentary abnormalities in the peripheral retina. In 2005, one of us published a clinical re-
port of an individual with aceruloplasminemia. In the present study, we examine the retinal histopathologic features of the same individual at age 60 years.

**TISSUE PREPARATION**

Eyes were obtained from a 60-year-old male donor with aceruloplasminemia after a 7-hour postmortem interval, in accordance with the tenets of the Declaration of Helsinki. At the time of the previous study, the donor had iron deficiency anemia, long-standing diabetes mellitus, cardiomyopathy, iron accumulation in the liver, dementia, and macular degeneration characterized by drusen and areas of retinal pigment epithelium (RPE) depigmentation. Subsequent to the publication of the report, he had experienced many hypoglycemic episodes, had developed renal failure, had been hospitalized, had developed aspiration pneumonia and sepsis, and had died. For the present study, samples from the anterior segment; retina, choroid, and sclera; and optic nerve of his right eye were used. As a normal control, an eye from a 60-year-old male donor with a 7-hour postmortem interval and no history of retinal disease was obtained from the Lions Eye Bank. A control eye with AMD was obtained from an 86-year-old woman with a history of AMD after a postmortem interval of 20 hours from the Complications of Age-Related Macular Degeneration Prevention Trial Eye Donor Program in collaboration with the Foundation Fighting Blindness. All eyes were fixed in formalin, embedded in paraffin, and sectioned at 7-µm thickness.

**PERIODIC ACID–SCHIFF STAINING**

Paraffin sections from the ocular regions indicated herein were stained using a periodic acid–Schiff stain kit (Polysciences, Inc, Warrington, Pennsylvania) as suggested by the manufacturer. Images were acquired on a Nikon Eclipse TE300 camera (Nikon Corporation, Tokyo, Japan) using Image-Pro Plus software, version 6.1 (Media Cybernetics, Inc, Bethesda, Maryland). Histopathologic analysis was performed as previously described.

**PERLS PRUSSIAN BLUE STAINING FOR IRON**

Paraffin sections were bleached with 0.2% potassium permanganate and 0.50% oxalic acid. Then, they were stained with 3% potassium ferrocyanide and 5% hydrochloride for 30 minutes. Paraffin sections from the ocular regions indicated herein were stained using periodic acid–Schiff stain kit (Polysciences, Inc, Warrington, Pennsylvania) as suggested by the manufacturer. Images were acquired on a Nikon Eclipse TE300 camera (Nikon Corporation, Tokyo, Japan) using Image-Pro Plus software, version 6.1 (Media Cybernetics, Inc, Bethesda, Maryland). Histopathologic analysis was performed as previously described.

**IMMUNOHISTOCHEMISTRY**

Paraffin sections were stained using the Vectastain ABC Alkaline Phosphatase kit for Rabbit and Mouse IgG and Vector BCIP-NBT kits (Vector Laboratories, Inc, Burlingame, California). Primary antibodies for l-ferritin (F-17; gift of Paolo Arosio, PhD, and Paolo Santambrogio, PhD), C5b-9 (M0777; Dako, Carpinteria, California), and vitronectin (MAB1945; Millipore, Billerica, Massachusetts) were used at 1:100 dilutions.

**ELECTRON MICROSCOPY**

The temporal macula was postfixed with osmium tetroxide, dehydrated, and embedded in EMBed-812 (Electron Microscopy Sciences, Hatfield, Pennsylvania). Thin sections were stained and examined with a JEOL1010 transmission electron microscope (JEOL Ltd, Tokyo, Japan). Images were acquired with Advanced Microscopy Techniques Image Capture software (Advanced Microscopy Techniques, Corp, Woburn, Massachusetts) and were rotated and cropped with Adobe Photoshop (Adobe Systems Incorporated, San Jose, California).

**RESULTS**

To determine whether the aceruloplasminemic eye had any unique pathologic manifestations, it was examined at the gross, light microscopic, and electron microscopic levels. At the gross level, the phakic right eye had no abnormalities other than macular degeneration, consistent with the results of the previous clinical report. At the light microscopic level, the cornea, iris, ciliary body, and optic nerve had normal morphologic findings (data not shown). The peripheral and macular neural retina had normal findings (Figure 1A and B), but the RPE cells and Bruch’s membrane contained many pathologic manifestations on all sections examined, similar to those previously described in AMD. Nodular drusen,16,19-21 were present in the macular, although they were much more common in the periphery (Figure 1C). The periphery also had many diffuse drusen,16,19,20 (Figure 1D), but the macula had none. The macula had subretinal drusenoid deposits22,23 (Figure 1E); the periphery did not. The macula and periphery had RPE cells with inclu-usions16,21 (Figure 1F) and RPE cells that had extruded from their normal layer16,24 (Figure 1G). The macula had several areas of RPE hypertrophy16 (Figure 1H) and atrophy16 (Figure 1I); the periphery did not. The macula and the periphery had areas of depigmented RPE cells that lacked the brown color of their neighbors and instead stained purple (Figure 1J and K); normal control and AMD RPE cells lacked this color variation (Figure 1L and M).

At the electron microscopic level, drusen and basal linear deposits resembled deposits previously described in AMD.25-27 (Figure 2A-E). Also, similar to AMD, aceruloplasminemic drusen had evidence of complement activation28-31 (Figure 3A-D).

To determine whether iron had accumulated in any cells, the Perls stain was performed. Strong staining was seen in the RPE of the periphery and the macula (Figure 4B, D, F, H, and J), but the periphery did not have as many Perls-positive cells. Hypertrophic RPE cells...
though the aceruloplasminemic macula had a strong ferritin staining than the periphery (Figure 5G-N). Al-

gage-related macular degeneration (AMD).

were always Perls positive (Figure 4J). The neural retina, choroid, optic nerve, ciliary body, iris, trabecular meshwork, and cornea were Perls negative (data not shown). Of interest, many RPE cells that subsequently had a strong Perls-positive label could never be fully bleached but retained a golden color, presumably from the high iron ac-
cumulation (Figure 4E-J).

To determine whether the neural retina and Perls-
negative RPE cells had increased iron levels compared with control cells, immunohistochemistry for the iron storage protein L-ferritin was performed. The neural retina (Figure 5F) and peripheral and macular RPE cells had strong L-ferritin staining, more than that of the normal control retina (Figure 5G-N). In control and aceruloplasminemic RPE cells, the macula had stronger L-ferritin staining than the periphery (Figure 5G-N). Al-

ough the aceruloplasminemic macula had a strong L-ferritin label, the golden cells with large iron deposits did not stain strongly with L-ferritin, suggesting that iron in them may not be stored as ferritin (Figure 5L and N).

To determine whether increased iron levels affected the normal RPE cytoarchitecture, cells were examined by electron microscopy. The aceruloplasminemic RPE mito-

chondria had abnormal electron-dense inclusions (Figure 6C and F). Normal RPE cells lacked these inclusions (Figure 6A and D), but RPE cells with AMD had occasional inclusions (Figure 6B and E). Such inclusions have not been previously described in RPE mito-

chondria, to our knowledge. Analysis via SIMS detected no increase in iron in the inclusions above the background tissue level, suggesting that the inclusions are areas of damaged lipids or proteins and not areas of significant iron deposits (data not shown).
Electron microscopic examination of unstained sections revealed 2 major types of RPE cells: melanosome-poor cells that had many electron-dense particles (Figure 7A, arrows) and melanosome-rich cells that lacked the electron-dense particles (Figure 7A, arrowheads). Such particles were never seen in eyes with AMD and normal control eyes (data not shown) and have not been previously reported, to our knowledge. Stained sections revealed melanosome-rich, melanosome-poor, and a few intermediate cells, suggesting a possible progression from the melanosome-rich to the melanosome-poor phenotype over time (Figure 7B-D).

In addition to melanosomes, the melanosome-rich cells contained lipofuscin and complex granules; all these structures are seen in AMD and with aging (Figure 7E). For 16 melanosome-rich RPE cells, the mean (SD) numbers of melanosomes, lipofuscin, and complex granules per cellular cross-section were 20.0 (7.8), 35.0 (12.0), and 11.0 (6.1), respectively.

In the melanosome-rich cells, a progression from a normal amount of melanosomes to large aggregates of melanosomes in lipofuscinoid material was observed (Figure 7F-L). The melanosomes in these aggregates could be intact (Figure 7F) or in different states of degradation (Figure 7K and L). Intermediate cells contained many more aggregates of degrading melanosomes than the melanosome-rich cells. In most cases, intact melanosomes no longer could be distinguished (Figure 7M). Aggregates with degrading melanosomes were not observed in normal control eyes or in control eyes with AMD (data not shown).

Melanosome-poor RPE cells lacked melanosomes, lipofuscin, and complex granules. For 16 melanosome-poor RPE cells, the mean (SD) number of melanosomes, lipofuscin, and complex granules per cell profile were 1.8 (2.8), 1.0 (1.5), and 0.8 (1.4), respectively. Melanosomes and complex granules were never seen outside the RPE in the pre-RPE or sub-RPE spaces, suggesting that melanosomes had not been extruded but had been degraded. Melanosome-poor RPE cells had many electron-dense, irregularly shaped membrane-bound granules that resembled siderosomes (Figure 7N-P). Such granules were never seen in normal RPE cells or RPE cells with AMD (data not shown) and have not been previously described in the RPE, to our knowledge.

To determine the subcellular location of iron in the RPE, unstained sections were examined by electron microscopy and SIMS. Complex granules from melanosome-rich aceruloplasminemic RPE cells had thin, grainy electron-dense borders (Figure 8A, arrowheads), but similar granules from normal RPE cells and RPE cells with AMD lacked these borders (data not shown). Melanosome-rich aceruloplasminemic RPE cells also had vesicles with granular electron-dense material (Figure 8A, arrow), but normal cells and those with AMD did not (data not shown). Analysis via SIMS demonstrated that the grainy borders on the outer edges of the complex and lipofuscin granules were iron, as was the material in the vesicles (Figure 8A [arrow], D, and E). Melanosomes contained some iron and high sulfur levels (Figure 8C-F).

The melanosome-poor cells had only siderosomes (Figure 8G) that contained very high amounts of iron, phosphorus, and sulfur (Figure 8G, J, and K). Compared with melanosome-rich cells, this iron content was much higher (Figure 8M and N).
We have presented for the first time, to our knowledge, the retinal histopathologic features of aceruloplasminemia and evidence for iron-mediated melanosome degradation. The aceruloplasminemic eye had many similarities to but also notable differences from eyes with AMD. Because this donor had died at age 60 years, it is possible that some of the AMD-like changes were the result of aging or AMD; however, disease was clinically evident from age 47 years, suggesting that most of our findings are the result of aceruloplasminemia.15

As in AMD, depigmentation, atrophy, hypertrophy, inclusions, and extrusion of the RPE were observed (Figure 1).16 These morphologic features of these conditions were similar to those seen in AMD; however, the extent of depigmentation was greater than in AMD. Depigmented cells are found mostly in the macula in AMD, but in aceruloplasminemia they were present in the macula and in the periphery.16

The aceruloplasminemic eye had drusen, basal linear deposits, and deposits in the subretinal space (Figure 1 and Figure 2).22,27 Although the morphologic findings regarding deposits were similar to AMD at the light microscopic and electron microscopic levels,16,23,26 the distribution differed from that which is typical of AMD. The aceruloplasminemic eye had many drusen in the periphery and fewer in the macula, but the opposite is true in AMD.16,20,31 The aceruloplasminemic eye also had a greater number of subretinal drusenoid deposits in the macula than is typical in AMD.32

In AMD, drusen contain activated complement. The terminal component of the complement cascade, C5b-9 (membrane attack complex), is present in most AMD nodular and diffuse drusen in a mottled pattern.29,36 The aceruloplasminemic drusen had positive results for C5b-9 and had the expected mottled pattern; however, not all drusen were labeled (data not shown). Aceruloplasminemic diffuse drusen had more frequent C5b-9 staining than did nodular drusen, but in AMD staining is similar in both types.36

Vitronectin, a plasma protein that inhibits the formation of the membrane attack complex, is present in high levels in all AMD drusen28,30 and is also seen in occasional RPE cells.30 Similar to AMD, vitronectin strongly labeled all aceruloplasminemic drusen and occasional RPE cells (Figure 3). Subretinal drusenoid deposits had negative results for C5b-9 and vitronectin (Figure 3).

Eyes with AMD have increased iron levels in the RPE compared with normal controls, as detected by enhanced Perls stain.35 As in AMD, aceruloplasminemic RPE cells had increased iron levels; however, the levels were so high that they could be detected by unenhanced Perls stain in all sections (Figure 4). Some aceruloplasmin-
emic RPE cells, especially in the macula, had so much iron that it was visible as a golden color that could not be bleached out (Figure 4). The aceruloplasminemic macula had more iron than the periphery (Figure 4); the same was true of the normal retina (Figure 5), suggesting that RPE cells of the macula and the periphery handle...
rather, neighboring cells that lacked the ferritin levels than normal RPE (Figure 5). Of interest, from the periphery and the macula had greater L-ferritin elevated (Figure 5). Also, aceruloplasminemic RPE cells and retinal astrocytes, increased iron levels in the aceruloplasminemic retina were expected.37,40 Indeed, iron distribution in maculas with AMD is more similar to that in the aceruloplasminemic periphery than in the aceruloplasminemic macula. Thus, the aceruloplasminemic periphery may better represent iron loading in AMD.

Aceruloplasminemic RPE cells had mitochondrial pathology (Figure 6). Most aceruloplasminemic mitochondria had at least 1 large focus of electron-dense material that was not iron but was likely damaged lipid and protein. The eye with AMD, which had a longer post-mortem interval than the aceruloplasminemic eye, had only occasional mitochondrial accumulations, which were much smaller than those in aceruloplasminemia. The presence of mitochondrial damage in aceruloplasminemia supports previous biochemical studies46-48 of aceruloplasminemic brains, which demonstrated mitochondrial dysfunction of respiratory chain complexes I and IV.

Melanosomes, lipofuscin, and complex granules of aceruloplasminemic RPE cells also differed from RPE cells with AMD and normal RPE cells (Figure 7 and Figure 8). A typical macular RPE cell from a normal 60-year-old eye has 17 to 20 lipofuscin, 11 to 13 melanin, and 9 to 14 complex granules per cell profile.17 The aceruloplasminemic RPE macular cells contained 2 types of cells, melanosome-rich and melanosome poor, both with abnormal numbers of granules. Melanosome-rich RPE cells had greater-than-expected numbers of lipofuscin granules and melanosomes per cell profile (35 and 20, respectively) but normal numbers of complex granules. The melanosome-poor cells, however, had significantly lower numbers of each granule type (approximately 1 of each per cell profile), suggesting that the melanosomes may have been degraded.

Melanosome degradation in the RPE is controversial,49 mainly because limited in vivo evidence has been published.46-51 Melanosome degradation has been described in fetal bovine RPE cells as the tapetum lucidum becomes amelanotic.50,51 As presented by Feeney-Burns and Mixon,50 the melanosomes in the tapetal RPE form aggregates of degrading melanosomes similar to the aggregates of degrading melanosomes that we observed in the aceruloplasminemic RPE (Figure 7). The mechanism of melanosome degradation in the tapetal RPE is not clear. Feeney-Burns and Mixon, along with the au-
thors of some contemporaneous studies,32,52,53 suggest that the presence of lysosomal enzymes in melanosomes implies melanosome degradation by lysosomes. However, it is now known that melanosomes are lysosome-related organelles that normally contain lysosomal enzymes and cannot be degraded easily.49,54,55

Melanosomes can, however, be photodegraded56,57 and degraded with hydrogen peroxide.59,60 Moreover, in vitro experiments have shown that the rate of synthetic melanosome-poor RPE cells has high sulfur levels in the siderosomes, suggesting that components of degraded melanosomes are present in the hemosiderin. Thus, the aceruloplasminemic RPE cells may be an in vivo example of what has been observed in vitro. In conclusion, our studies have shown that RPE cells are greatly affected by the loss of ceruloplasmin; they accumulate iron and develop an AMD-like condition with many unique features.

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